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Fig. 1.—Two high 16-inch plate mill (of the Bureau of Standards) in which the ingots were rolled

MANUFACTURE AND PROPERTIES OF STEEL PLATES CONTAINING ZIRCONIUM AND OTHER ELEMENTS

By George K. Burgess and Raymond W. Woodward

ABSTRACT

This paper describes the manufacture and certain physical properties obtained from steel plates produced from about 193 heats of steel containing in various combinations the following principal variable elements: Carbon, silicon, nickel, aluminum, titanium, zirconium, cerium, boron, copper, cobalt, uranium, molybdenum, chromium, and tungsten.

None of the steels presented any difficulties in rolling into plate except those containing boron. Boron forms a complex eutectic, probably that of an iron-carbon-boron compound with iron, which is fusible at the temperatures ordinarily used in rolling, but at slightly lower temperatures steel containing boron can be rolled successfully.

The usual mechanical and impact tests were carried out on all of the steels. It is shown that steel containing 0.40 to 0.50 per cent carbon, 1 to 1.50 per cent silicon, 3 to 3.25 per cent nickel, and 0.60 to 0.80 manganese and deoxidized with a simple deoxidizer, such as aluminum, can be produced having a tensile strength of approximately 300 000 lbs./in.² with excellent ductility and toughness. This type of steel is recommended for a structural material.

Although the same high properties are obtained in steels of the above composition with the aid of additional elements, it does not appear necessary in general, to resort to such additions of more costly alloying elements.

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I. INTRODUCTION

This investigation originated in the need of the ordnance departments of the Army and Navy for information regarding the effects on the ballistic properties of light armor plate of certain chemical elements, such as zirconium, on the one hand, and the effects of such elements as uranium in reducing erosion in guns, on the other hand. The account here given relates mainly to the efforts to produce armor plate of various compositions.

After conference with the representatives of the several establishments interested, a joint program was outlined according to which the Bureau of Mines was to produce and analyze ingots of the desired compositions, the Bureau of Standards to manufacture and heat treat plates, carry out physical tests, microexaminations and chemical analyses, and develop methods of chemical analysis when needed for the more unusual elements in steel, and the Navy Department was to carry out the ballistic tests.

The most urgent problem was the determination of the effect of zirconium on the properties of carbon steels and of nickelcarbon steels, and especially to differentiate between the effects of zirconium and silicon.

As the work progressed it was considered desirable to investigate the effects of other elements, and there were accordingly included steels containing titanium, aluminum, boron, molybdenum, cerium, cobalt, chromium, vanadium, tungsten, uranium, and copper.

In addition to the ingots furnished by Dr. Gillett, opportunity was given to examine plates of steel containing zirconium manufactured by an automobile manufacturer.

Although the results of the ballistic tests are not available for publication, an account of the mechanical properties and tests of this series of somewhat unusual steels is considered worthy of consideration. The nickel-silicon group appears to be of particular interest, as is also the fact brought out that zirconium does not appear to confer any especially advantageous properties to types of steel here studied, and, in fact, behaves very much like silicon, although for carbon steels the data are not sufficiently complete to warrant conclusions.

The investigation throws some additional light on the manner in which certain of the rarer elements enter into steel, and there were also developed new analytical methods for the determination of zirconium in steels and of several of the unusual elements in the presence of each other.

II. COMPOSITION OF INGOTS

The chemical composition of all ingots was determined from drillings taken from both the top and bottom of the ingots. From each of these sets of drillings an analysis was made for all the elements occurring in the steel by Dr. Gillett and his associates at Ithaca. Samples were also taken from the top and bottom crops at the Bureau of Standards and further analysis made for the aluminum, titanium, and zirconium content. Since the exact determination of this combination of elements presents some difficulty, the method used at the Bureau is included in the appendix.

The composition of the various ingots will be found in Tables 11 to 22, while Table 1 gives a list showing in which table the data for a given heat appears. The analytical values for carbon, silicon, manganese, nickel, and other alloying elements are as reported by Dr. Gillett. Those for aluminum, titanium, and zirconium are a weighted mean of the determinations made at the Bureau of Standards and by the Bureau of Mines. In case the top and bottom samples showed segregation to have occurred in the ingot the values for both top and bottom are given in the table. No determinations were made of the sulphur or phosphorus content except in the few cases noted, since the steels were made from Armco iron as a base, and it is believed that these steels will run below 0.035 per cent sulphur, except 1256 and 1257, in which it was intentionally raised, and below 0.015 per cent phosphorus.

¹ These determinations were made under the direction of Dr. G. E. F. Lundell.

² See also Lundell and Knowles, Jl. Ind. and Eng. Chem., 12, p. 562, 1920; The determination of zirconium in steel.

In addition all will contain about 0.04 per cent copper and probably a small amount of cobalt, carried in by the commercial nickel, in the steels containing nickel.

TABLE 1.—List Showing Tables in Which Composition and Mechanical Properties of Various Heats May be Found

	able Heat No. No.	Table No.	Heat No.	Table	Heat	Table
1101 1101 1	10.			No.	No.	No.
				110.		110.
1101 13 1163 1	1 1204	12	1243	14	1290	14
	1 1205	12	1244	22	1291	14
1103 13 1165 1	2 1206	12	1245	12 12	1292	14
	2 1207	20	1246	12	1293	14
1105 13 1167 1	2 1208	12	1247	14		!
1106 13 1168 1	2 1209	12	1248	14		parison
	2 1210	14	1249	14	st	els
	2 1211 2 1212	14 14	1250 1251	14 12		
	2 1212	14	1251	15		
					1 2	14 14
1113 12 1173 2	0 1214	12	1253	15	3	I8-21
	2 1215	12	1256	15	3 4 5	18
	4 1216	12	1257	15	5	19
1117 14 1176 1	4 1217	12	1258	15		1
1118 12 1177 2	2 1218	14	1259	15	6	18
1119 14 1178 2	2 1219	14	1260	15	7 8	14
1120 12 1180 1	3 1220	14	1261	17	8	14
1128 12 1181 1	3 1221	14	1263	17	9	18
1129 12 1182 1	3 1222	14	1264	17	10	14
1130 12 1183 1	3 1223	14	1267	17	11	14-18
					12	19
	3 1224	14	1268	15	13	12
1132 14 1185 1	3 1225	14	1269	11	14	14-21
	4 1226	12	1270	11	15	18-21
1134 14 1187 1	4 1227 4 1228	12	1271	20 15		
1135 18 1188 1	4 1228	22	1272	15	16	14
1136 18 1189 1	4 1229	22	1273	20	17	14
1138 14 1190 1	4 1230	14	1274	17	18	14
1144 14 1191 1	4 1231	14	1275	17	19	19
1145 14 1192 1	4 1232	14	1276	17	20	14
1146 14 1193 1	4 1233	14	1277	17	21	14
				1 10	21 22	21
1147 12 1194 1	4 1234	14	1278	17	23	20
1155 19 1195 1	4 1235	14	1279	16	24	14
1156 19 1196 1	4 1236	12	1280	16	25	14
	4 1237	12	1281	15-16		
1158 14 1198 1	1 1238	12	1282	16	26	14
1159 14 1199 1	1 1239	12	1283	16	27	14
1160 14 1200 1	1 1240	14	1285	16	28	12
	1 1241	14	1286	16		1.00
1162 14 1202 1	2 1242	12	1289	14		
		li .				

III. PREPARATION OF PLATES AND TEST PIECES

1. CROPPING AND ROLLING OF INGOTS

The ingots after having been made at the Bureau of Mines Experimental Station at Ithaca, N. Y., as described in a forth-coming paper of the Bureau of Mines, were shipped to the Bureau of Standards and there rolled into plates.

(a) DESCRIPTION OF INGOTS

The first ingots received were plain, round ingots without hot tops, cast large end up, the length being about 15 inches, top diameter about 3½ inches, and bottom diameter about 2½ inches.

This series, embracing Nos. 1101 to 1120, naturally contained a large pipe or cavity at the upper end, and a top crop of about 5 inches was necessary to remove physically unsound material.

Bottom crops on this series (except Nos. 1118 and 1120) had been taken previous to receipt at the Bureau of Standards. No crops were taken on ingot Nos. 1101–1104, 1109, 1111, and 1114 at the Bureau, as these ingots had been machined down on the ends and surface before shipment.

Ingots from No. 1128 to 1158 were also of the same general dimensions and form as the previous ones, but with the addition of a hot top of about 2½ inches diameter. Most of this top had been knocked off while the ingot was still hot, directly after stripping from the mold. The tops of these ingots were cropped just below the junction of the hot top with the body of the ingot, or slightly farther in a few cases, to insure sound material. The bottom crop was just sufficient to remove the rounded end of the ingot.

The remainder of the ingots were square in cross section, about 3 inches at the top and 2½ inches at the bottom. The length, exclusive of the hot tops, was about 21 inches. The average weight of these ingots was 41.5 pounds before cropping and 35 pounds when ready to roll. The length after cropping was about 18 inches.

Table 2 gives a summary of the weight of ingots and crops for the various types of the ingots and also the percentage available for rolling after having been cropped. The data show remarkably well the advantage to be gained by the use of a hot top, the available material without such means being about 65 per cent, while with a hot top the average was 84 per cent. This latter figure should be slightly reduced (possibly 5 per cent) to allow for a portion of the hot top having been knocked off at Ithaca. This corrected figure, however, agrees very well with similar data obtained on 4½-ton ingots and reported elsewhere, sepecially when the small size of the ingots used in this investigation is considered.

(b) ROLLING DATA

The ingots were rolled in a two-high 16-inch plate mill, a photograph of which is shown in Fig. 1. This mill is driven by a 150-horsepower 230-volt direct-current motor and is nonreversing.

²Steel Rails from Sink-Head and Ordinary Rail Ingots, by George K. Burgess, Bureau of Standards Technologic Paper, No. 178

The motor speed is variable from 250 to 1000 revolutions per minute, which with the reduction gear gives a roll speed of 20 to 80 revolutions per minute. For this work the roll speed was kept constant at the lowest value, corresponding to a peripheral velocity of approximately 83 feet per minute.

The heating of the ingots was by means of a gas-fired semimuffle furnace. Usually about 10 ingots were rolled at one heating, and they were charged into the cold furnace and brought to temperature with the furnace. With the exception of those steels containing boron, which are discussed later, the furnace temperature was maintained at from 1100 to 1150° C. The ingots were rolled until their temperature had fallen to about 850° C. This temperature was checked in many cases by means of an optical pyrometer. The ingots were then reheated and the operation repeated.

All of the ingots were first squared down to $2\frac{1}{2}$ inches square in four passes by turning the ingot through a 90° angle at alternate passes. This gave a maximum reduction of about 10 per cent per pass.

Ingots up to 1163 were then cross rolled until 6 to 7 inches wide and then rolled lengthwise until ½ inch thick. The average size of finished plate was about 28 by 6½ by ½ inch. A total of about 40 passes was required, with about 5 per cent reduction per pass. The other ingots were entirely cross rolled after squaring, producing plates of various sizes, usually about 20 by 13 by ½ inch (or ¾ inch).

It is probable that the percentage reductions per pass could be considerably increased, but since the rolling properties of the majority of the ingots were unknown it was deemed advisable to have a considerable margin of safety. A portion of those ingots which were rolled into long narrow plates (1101 to 1162) was cut off and rerolled to 3/8 inch thickness.

TABLE 2.—Average Ingot Weights

Ingot Nos.	Original weight	Тор сгор	Bottom crop	Cropped ingot	Available for rolling	Loss in rolling
7	Pounds	Pounds	Pounds	Pounds	Per cent	Per cent
1101-1120	34. 5	12.0		21. 2	64.9	3, 1
1128-1158	34. 5	5.9	1.5	27.1	79.8	2.3
1159-1293	41.5	4.8	1.8	35.0	84. 2	1.9
Average of all	40. 5	5.3	1.7	32. 7	82. 4	2.0

(c) DISPOSITION OF MATERIAL

The plates from ingots 1101 to 1162 were sawed similar to sketch of Fig. 2, which represents plate No. 1129, although typical of all. AB and GF were plates for ballistic tests, BC and CD

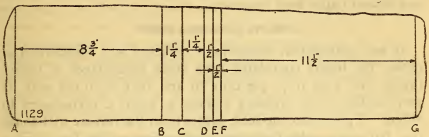


Fig. 2.—Disposition of material from plates rolled from small ingots

each about r inch wide, were for tensile specimens, EF for microscopic examination and thermal analyses, while the remaining small pieces were held in reserve. The lengths of the two plates were so adjusted that the rerolled portion would be the same length as the unrolled portion.

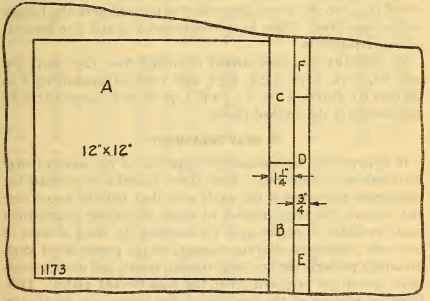


Fig. 3.—Disposition of material from plates rolled from large ingots

From the later plates (1163 to 1293) a single ballistic test piece was cut 12 by 12 inches, or as large as possible, if the ingot was too small to permit of this dimension. The other test pieces were taken then as shown by the typical diagram for plate No. 1173 in Fig. 3. A is the ballistic plate, B and C tensile specimens,

D for impact sample, and E and F for microscopic examination and thermal analyses, respectively. Any plates which were not perfectly flat were straightened at a temperature of about 800° C by means of a hydraulic forging press. The plates were not pickled until after heat treatment.

(d) INGOTS CONTAINING BORON

As was anticipated, considerable difficulty was experienced in rolling the ingots containing even small percentages of boron. Ingots Nos. 1254 (0.73 per cent B) and 1255 (0.30 per cent B) were heated in the ordinary manner to 1100° C preparatory to rolling. Upon removing these ingots from the furnace with tongs they fell apart under their own overhanging weight, furnishing a striking example of hot-shortness. No. 1262, containing 0.46 per cent boron, was heated to only 960° C and likewise broke after partial rolling. Nos. 1261, 1263, and 1264 were similarly heated and were rolled satisfactorily, although containing 0.71, 0.23, and 0.46 per cent boron, respectively. Nos. 1265 and 1266 were partially rolled, but broke up under the rolls and, in fact, were so hard to roll that two of the coupling collars of the mill were also broken at the same time. These ingots contained 0.23 and 0.46 per cent boron, respectively.

No difficulty was encountered in rolling Nos. 1267 (0.44 per cent B), 1274, 1275, 1276, 1277, and 1278 (all containing 0.10 per cent B), although Nos. 1274 and 1276 showed numerous cracks and fissures in the finished plates.

2. HEAT TREATMENT

In determining the mechanical properties of the series of steel two needs were considered. First, it was desirable to correlate the mechanical properties of the steels with their ballistic properties; and, second, the large number of steels of varying composition made available an opportunity for studying the effect of some of the more uncommon alloying elements on the properties of steel. To satisfy properly the first requirement, tensile and impact specimens should be prepared from the heat-treated plates. This, however, because of the hardness of the plates, was practically impossible to carry out without what seemed to be unwarranted expense and time. For the second requirement it would be preferable to make tests on a series of specimens from each composition drawn back to various temperatures after quenching from the proper temperature. This, again, would have required con-

siderable additional work; and, moreover, there was not enough material to make a complete survey of the entire tempering range.

A compromise was accordingly effected whereby tensile tests were made on normalized and heat-treated specimens cut from the plates before heat treatment, as noted in Section III, 1 (c). The heat treatments were similar for all compositions of the series. That is, after normalizing and quenching in oil each from a temperature 30° C above the end of the upper critical range the specimens were drawn back at a temperature 175° C for three hours, this being the same treatment given the plates for ballistic testing.

TABLE 3.-Normalizing and Hardening Temperatures

No.	Temp.	No.	Temp.	No.	Temp.	No.	Temp.	No.	Temp.
1101	840	1159	780	1195	840	1231	800	1271	780
1102	860	1160	780	1196	860	1232	780	1272	820
1102	825	1161	840	1197	860	1232	780	1273	820
1103	840	1162	780	1197	900	1233	840	1274	820
1104	810		860	1198	900				
1105	810	1163	800	1199	900	1235	860	1275	840
1106	860	1164	860	1200	820	1236	840	1276	760
1107	860	1165	820	1201	840	1237	840	1277	800
1109	800	1166	860	1202	860	1238	800	1278	850
1111	770	1167	780	1204	840	1239	800	1279	760
1112	790	1168	840	1205	820	1240	800	1280	780
1113	770	1169	820	1206	820	1241	780	1281	780
1114	770	1170	780	1207	800	1242	800	1282	800
1115	780	1171	860	1208	800	1243	800	1283	780
1117	800	1172	860	1209	800	1244	820	1285	820
1118	785	1173	840	1210	860	1245	820	1286	800
1119	810	1174	840	1211	860	1246	800	1289	800
1120	820	1175	840	1212	800	1247	840	1290	820
1128	820	1176	820	1213	800	1248	880	1291	780
1129	840	1177	860	1214	820	1249	800	1292	780
1130	820	1178	860	1215	800	1250	920	1292	800
1131	830	1180	860	1216	820	1251	820		
1132	830	1181	860	1217	800	1252	800		
1133	820	1182	860	1218	820	1253	805		
1134	820	1183	860	1219	820	1256	840		
1135	780	1184	900	1220	800	1257	840		
1136	780	1185	900	1221	820	1258	780		
1138	805	1186	900	1222	780	1259	780		1000
1144	820	1187	900	1223	780	1260	820		
1145	820	1188	900	1224	820	1261	880		
1145	780	1189	900	1225	820	1263	880		
1147	790	1190	860	1226	840	1264	820		
1147 1155	805	1191	860	1227	840	1267	880		
1156	805	1192	860	1228	800	1268	840		
1157	810	1193	860	1229	820	1269	840		
1158	780	1194	840	1230	760	1270	840		

The normalizing and hardening treatments for the smalltest specimens were carried out in a small electric muffle furnace; the specimens were placed in the cold furnace, brought to the desired temperature with the furnace, and held at that temperature for 15 minutes. The normalized specimens were allowed to cool in the air, and the specimens to be hardened were quenched in the oil at room temperature. In Table 3 are given the common normalizing and quenching temperatures. For tempering, the specimens were heated in an oil bath to 175° for three hours and allowed to cool in the air after removing from the bath.

The duplicate tensile specimens from each plate were, in general, normalized at the same time, and then one of them hardened and tempered and the other kept in the normalized condition.

The ballistic plates were heat treated in a manner similar to that used for the test specimens, except that larger furnace units were required and the plates were held at the desired temperatures for 45 minutes before withdrawing from the furnace.

IV. PROPERTIES OF THE MATERIAL

1. CRITICAL RANGES

In order to prescribe properly the heat treatment of the various steels, the critical ranges of several of them were determined, particularly those containing the more unusual elements, such as zirconium, boron, cerium, copper, and large amounts of silicon. Inverse rate curves were obtained on samples of 1.5 to 2.0 grams mass by means of a modified Rosenhain furnace.4

The temperature measurements were taken with a platinum, 90-platinum 10-rhodium thermocouple. The rate of heating and cooling was approximately 0.20° C per second, while the maximum temperature to which the specimen was carried varied between 840 and 920° C.

Table 4 gives the results obtained, together with the composition of the materials investigated. Although the end of the Acas transformation is all that is needed to determine the proper heat treatment, the other values are included as a matter of interest. For comparison there are also shown in the table data for a plain carbon and a 3 per cent nickel steel, both of which contain the other elements within commercial limits.

⁴ Scott and Freeman, Bull. Am. Inst. of Min. and Met. Eng., No. 152, p. 1429; August, 1919. Also Bureau of Standards Scientific Paper, No. 348.

654 572

672

681 617

729

743

802

783

748

732

726

TABLE 4.—Critical Ranges of Representative Material

[All temperatures in ° C]

43							
t	-	End	668 668 664 670 671	680 668 577 588 578	572 594 588	590 668 675 547	
	Arı	Max.	690 677 676 683 683	690 686 605 613 613	611 605 426 617	613 684 690 603	
		Beg.	695 684 682 688 690	696 692 615	616 477 620	622 699 694 611	
	Ar ₃₋₂	Max.	716 755 751 710 731	775 718 636	641	685 722 714 616	
	Ψ	Beg.	745 784 773 730 755	798 749 659 646 658	643 671 658	701 738 734 628	
	Ac ₂ -3	End	780 842 ? 820 ? 778 828	843 808 739 758 738	736 778 782	817 793 785 725	
	AC.	Max.	7683	7977	776	795 777 778 714	
		End	778 770 756 759 759	766	746 751 741	745	
-	Acı	Max.	759 754 738 747	753 753 703 725 716	705 733 720 725	739 737 738 697	
		Beg.	755 750 732 741	745 748 697 720 711	701 719 719	732 734 682	FEELS
ner Commence of the Commence o		Other elements			Mo 0.78 B .30	Cu. 62	COMPARISON STEELS
		ZC	0.10	.20 .11 .03 .08	.30		OMPA
		TI	0.0 . 06 . 03 . 01	839.5.88	.03		
	Composition	Aı	0.09 .03 .01 .07	.03	.01	.03	
the objection of the same	Com	Ni		3.15	3.3.3 3.3.3 2.3.3 2.90	2. 80	
		Mn	0.80 .65 .58 .75	.50 .78 .59 .76	.65 .73 .83 .75	. 69 . 69 . 69 . 69	
		·IS	1.15 1.15 . 66 . 54 . 44	. 73 . 85 . 27 1. 10 1. 15	. 52 1. 65 1. 45 1. 00	1.30 .27 .33 .23	
		υ	0.51 .36 .39 .56	.37 .44 .46 .55	38	.16	
		No.	1101 1102 1104 1105	1107 1109 1111 1112 1113	1114 1133 1135 1255	1262 1272 1274 1279	

a These values are taken from Bureau of Standards Scientific Papers, No. 376.

2.90

0.75

0.22

0.40

C 24.

By comparing C 24, 1104, and 1102, containing, respectively, 0.22, 0.66, and 1.15 per cent silicon, it will be noticed that silicon progressively raises the Ac ranges and the Ar_{3-2} range. The Ar_1 point is also raised, but not to so great an extent as the other values. This effect is also the same in the presence of nickel, as shown by Nos. 1111 and 1112.

Zirconium has about the same effect on the critical ranges as silicon, as will be observed from an inspection of the several data.

The effect of cerium is apparently rather small and irregular, as will be seen from Nos. C24 and 1272. The Ac_1 and Ar_1 points are raised somewhat and the Ac_{2-3} and Ar_{3-2} points decreased by about a similar amount.

Copper evidently produces the same result upon the critical ranges as an equal amount of nickel, No. 1279 having values that would be expected for a 3 per cent nickel steel with 0.58 per cent carbon.

In specimen No. 1135, containing 0.78 per cent of molybdenum, the Ac₁ range is about normal for a steel of similar composition but without the molybdenum, whereas the Ar₁ range has been considerably depressed, as has also been observed by other investigators.

From Nos. 1255, 1262, and 1264, containing boron in amounts from 0.06 to 0.49 per cent, and by comparison with Nos. C 24 and 1133, keeping in mind the variations in composition, it appears that boron raises all of the ranges somewhat.

2. MICROSTRUCTURE

(By S. Epstein)

It must be assumed that if zirconium or any of the other elements that were used are to have any virtue as additions to light armorplate steels they should exert some noticeable effect on the microstructure of these steels at least in some stage of the heat-treated state, if not in the annealed condition. Also, if any true comparisons are to be drawn between steels of different chemical compositions from tests of their mechanical properties, it is plainly essential to know that the tested specimens have all been treated alike, or, if there are differences in the treatments, to know where they occur. It was the object, therefore, in the microscopic examination first to determine the rôle played by the zirconium, as well as the other addition elements in the steels, and, second, to examine the plates and test specimens for variations in soundness and heat treatment.

(a) ZIRCONIUM, TITANIUM, ALUMINUM

A steel containing zirconium can at once be recognized under the microscope by the presence of bright-yellow square inclusions not plainly visible at magnifications lower than 500 diameters (see Fig. 4). They are not affected by the ordinary alcoholic nitric or picric acid etching solutions, but retain their lustrous yellow color. In the steels examined these yellow inclusions were generally associated with orange-pink square inclusions and irregularly shaped purplish-gray ones. By a comparison of the chemical compositions of the steels the orange-pink ones were traced to the presence of titanium, while the purlpish-gray ones are probably due to alumium. However, this last point could not be established as positively as with the yellow and orange-pink inclusions (see Fig. 5). All of the inclusions manifested a tendency to form groups of tiny segregates, which when rolled, flattened out to thin plate-like streaks (see Fig. 6). These plates could readily be seen with the naked eye in the polished and etched specimens as yellow streaks. They could be noticed, also, in the fractures of some of the tested tension bars and impact specimens as laminations. In the specimens containing zirconium and titanium in which cracks were found the inclusions were most numerous very close to the cracks. Very few of the inclusions were found outside the segregates and streaks or away from the cracks.

Except for the bright-yellow square inclusions, which persisted throughout the working of the steel, the presence of zirconium was not found to affect the microstructure of the steels in any respect. Although the steels examined were regarded for the most part as alloy steels, the majority of the series under the microscope looked like plain carbon steels (see Fig. 7). In the air-cooled specimens it was considered that if any martensite or troostite were found it must be attributed to a self-hardening property or retardation of the Ar, transformation produced by the alloying elements. Otherwise, the structure would consist of granular pearlite and sorbite. Many of the other steels, to be referred to later (see Fig. 8), did show some martensite and troostite in the normalized specimens; but in no case was the presence of martensite or troostite in air-cooled specimens found to be due to zirconium. It may be that not all of the zirconium goes to form the characteristic yellow squares and that some of it goes into solution in the steel; but in that case it does not have a marked effect on the microstructure. In this

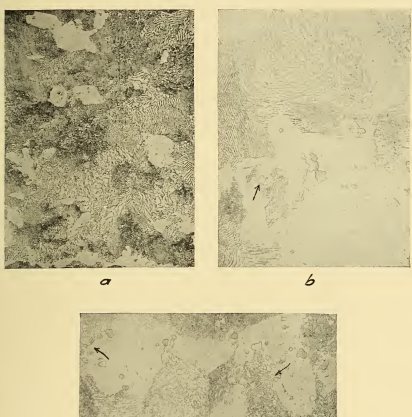
respect it may be likened to silicon, which goes into solution but the presence of which in small amounts can not be detected under the microscope. Thermal analyses seem to indicate that part at least goes into solution. Titanium and aluminum also formed characteristic inclusions, but otherwise gave no sign of their presence under the microscope.

The yellow square inclusions are very hard, but because they make up such a small proportion of the mass of the steel it can scarcely be conceived that they can exert a very great influence on its mechanical properties. The fact that they are associated in the form of segregates is a disadvantage, especially in an armorplate steel. Wherever cracks were found in steels containing zirconium the yellow inclusions were most numerous near the cracks. and the conclusion that the cracks are in some way associated with the plates of inclusions appears to be warranted. The yellow inclusions of zirconium are very similar to the orange-pink ones of titanium, and with regard to the tendency of the former to segregate and form its negative effect on the microstructure it may also be compared to titanium, which is regarded as a scavenger and not a true alloying element. In general, zirconium, titanium, and aluminum may all be put in one class. They appear to act primarily as scavengers, and when they are not removed as part of the slag are present in the steel in the form of inclusions. They may go into solution in the steel, but in that case their presence can not be detected in the microscope and their effect would appear to be slight or negligible. In the form of inclusions they can not do much good, and if these are segregated and rolled out into thin plate-like streaks they may be detrimental.

(b) OTHER ALLOYING ELEMENTS

In contrast to zirconium, titanium, and aluminum, the other additions to the steels may be regarded as true alloying elements. Carbon, silicon, manganese, and nickel were not considered as special additions in this respect, and while their presence in the course of the examination was always noted they were not under particular observation. The other alloying elements may be grouped as follows: (1) Chromium, tungsten, vanadium, molybdenum; (2) cerium, uranium; (3) copper; (4) boron.

The first four—chromium, tungsten, vanadium, and molybdenum—go into solution and produce a martensitic pattern in the air-cooled specimens (see Fig. 8, a, b, c, d). Cerium and uranium go into solution and produce a martensitic pattern in the



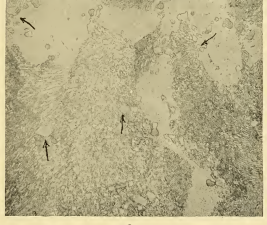


Fig. 4.—Inclusions in ingot 1109 as cast containing 0.11 per cent zirconium. Etching, 2 per cent nitric acid

(a) The bright yellow inclusions can not be distinguished at this magnification. X100 (b) This spot shows the average number of inclusions which are indicated by the arrows. They are bright yellow in color. X500 (c) A segregate of the bright yellow square inclusions is here shown. Most of these inclusions are segregated in this way. X500

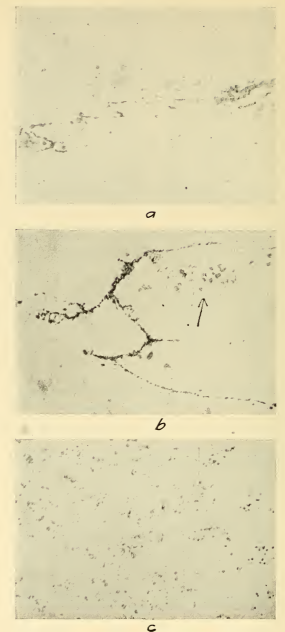


Fig. 5.—Different types of inclusions found in the steels. Not etched. Magnification

(a) Steel 1158. Al, 0.173 per cent; Ti, 0.028 per cent; Zr, 0.22 per cent. Of the square inclusions some are bright yellow and some are orange-pink. The yellow inclusions are due to zirconium, the orange-pink to titanium. The circular inclusions arranged in a threadlike continuity are purplish gray in color and may be due to aluminum (b) Steel 1176. Al, 0.25 per cent; Ti, 0.09 per cent; Zr, 0.11 per cent. A threadlike streak of the gray similar to (a) is to be seen. In the cluster of square inclusions indicated by the arrow the larger light-colored ones are bright yellow, the smaller darker ones are orange-pink
(c) Steel 1213. Ti, 0.04 per cent; Zr, 0.34 per cent. This shows a segregate of small yellow and orange-pink inclusions. Most of these inclusions are found in these segregates, very few outside

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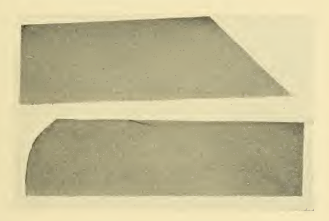


Fig. 6.—Segregates of Ti and Zr inclusions rolled out into tin plate-like streaks

Steel 1211. Butt—Ti, 0.07 per cent; Zr, 0.77 per cent. Top—Ti, 0.06 per cent; Zr, 0.92 per cent. The thin plates appear as yellow streaks easily visible to the naked eye in the polished and etched specimens. In the fractures of the impact and tension bars they look like laminations. Not etched. X2

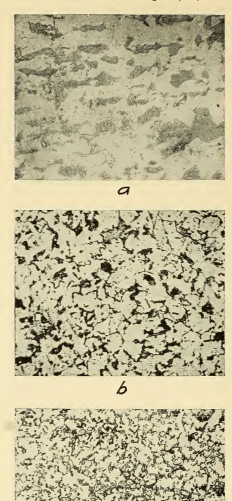


Fig. 7.—Microstructure of air-cooled specimens of steel containing Ti and Zr. Etching, 2 per cent nitric acid; magnification, ×500

(a) Steel 1185. C, 0.26 per cent; Ni, —; Al, 0.021 per cent; Zr, 0.30 per cent. The structure is granular pearlite and ferrite
(b) Steel 1186. C, 0.26 per cent; Ni, 2 per cent; Al, 0.013 per cent; Ti, 0.04 per cent; Zr, 0.24 per cent. The structure is again granular pearlite and ferrite. The structure is not essentially different from that of a steel containing a similar percentage of nickel but no other additions

similar percentage of nicker but no other additions
(c) Steel 1225. C, 0.27 per cent; Ni, 3.04 per cent; Ti, 0.22 per cent; Zr, 0.34 per cent. The structure is granular pearlite and ferrite. In no case was the presence of Ti or Zr found to produce a martensitic pattern

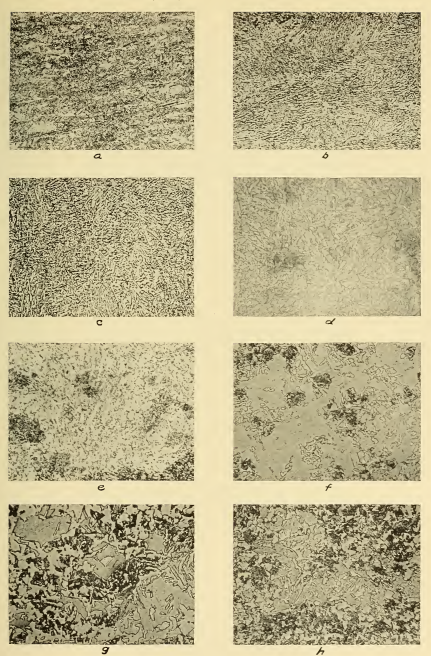
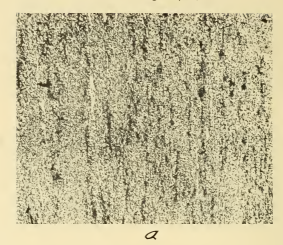


Fig. 8.—Microstructure of air-cooled specimens containing other alloying elements. Etching, 2 per cent nitric acid; magnification, × 500

- (a) Steel 1155. C, 0.38 per cent; Ni, 3.60 per cent; Cr, 1.14 per cent. The angular structure is typical of air-cooled nickel chromium steels
 (b) Steel 1178. C, 0.32 per cent; Ni, 3.5 per cent; Cr, 2 per cent; W, 0.90 per cent. The angular structure is characteristic
 (c) Steel 1207. C, 0.60 per cent; Ni, 3.5 per cent; Mo, 0.78 per cent
 (d) Steel 1135. C, 0.39 per cent; Ni, 3.5 per cent; Mo, 0.78 per cent
 (e) Steel 1258. C, 0.39 per cent; Ni, 3.65 per cent; Ce, 1.35 per cent. This steel displays a characteristic martensitic structure, due doubtless to the presence of the cerium
 (f) Steel 1228. C, 0.63 per cent; Ni, 3.01 per cent; N, 0.52 per cent. The uranium has produced a
 martensitic pattern
 (g) Steel 1226. C, 0.40 per cent; Si 1.61 per cent; M, 0.00 per cent; Ni, 3.04 per cent. The combination

- (g) Steel 1226. C, 0.40 per cent; Si 1.61 per cent; M, 0.90 per cent; Ni, 3.04 per cent. The combination of high Si, Mn, and Ni has resulted in a martensitic pattern (h) Steel 1282. C, 0.45 per cent; Si, 1.10 per cent; Mn, 0.84 per cent; Ni, 1.92 per cent; Cu, 1.35 per cent. The high percentage of copper had produced some martensite



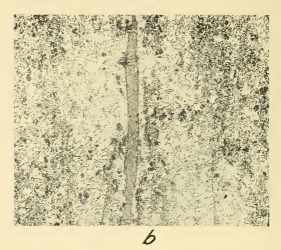


Fig. 9.—Inclusions in cerium and uranium steels. Etching, 2 per cent nitric acid

(a) Steel 1252. Air-cooled, Ce, 0.55 per cent. A large number of inclusions were segregated in streaks in this specimen. These streaks could be plainly seen with the naked eye. The shape and color of the inclusions can not be distinguished at the magnification of this micrograph, but at 500 diameters most of them appear as circular gray inclusions, while some are orange with gray markings inside the circumferences. The latter inclusions were found only in the cerium steels. X50 (b) Steel 1228. Air-cooled uranium, 0.52 per cent. A large number of inclusions were segregated in this spot, together with the long slag inclusion shown here. At higher magnification it can be seen that the inclusions are circular and of a deep blue color. The typical angular structure resulting from the presence of uranium is especially noticeable in this segregated area. X100





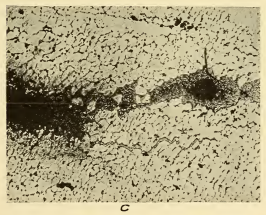


Fig. 10.—Characteristic structure of boron steel

- (a) Steel 1262, containing 0.49 per cent B, which broke in the rolls. This shows the eutectic network. Not etched. ×100
 (b) The eutectic etches dark with sodium picrate. This also shows where some of the eutectic has ccalesced into the circular particles. Etching, sodium picrate. ×50 (c) The eutectic is fusible at the temperature ordinarily used in rolling. The cracks in the broken ingot all followed the network of the eutectic. This micrograph shows where a larger mass of eutectic has located a crack. Etching, sodium picrate. ×50

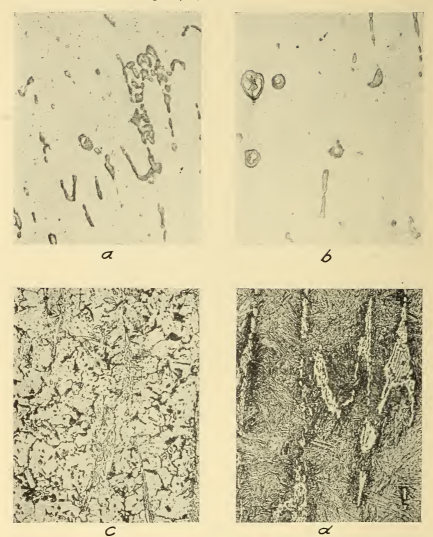
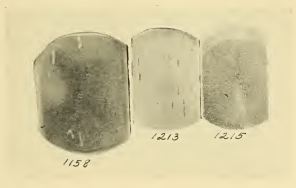


Fig. 11.—Rolled and heat-treated steels containing boron, × 500

- (a) Steel 1275. B, 0.08 per cent. This shows particles of the boron compound in the rolled and normalized specimen. Etching, sodium picrate
 (b) Same steel as (a). This shows the particles of the boron compound in the quenched specimen. Etching, sodium picrate
 (c) Steel 1263. B, 0.30 per cent. This is the air-cooled specimen. The white sharply outlined particles are the boron compound. Etching, 2 per cent nitric acid
 (d) Same steel as (c), quenched. There are no definite circular and elongated particles, but a eutectic structure is present. This may be due to the fact that the specimen was quenched from a temperature high enough to allow the eutectic to form again. Etching, 2 per cent nitric acid



a

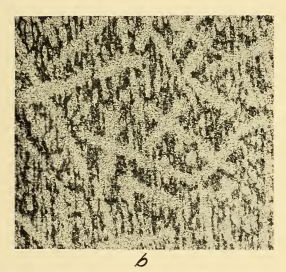
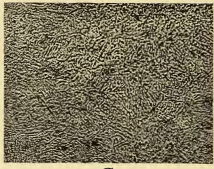


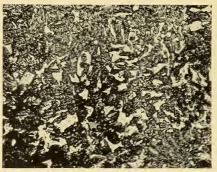
Fig. 12.—Types of flaws existing in some of the steels. Etching, 2 per cent nitric acid

(a) Steel 1158, H. T., cracked. Steel 1213, H. T., yellow streaks of titanium and zirconium inclusions. Steel 1215, H. T., dendritic. These specimens were taken from the shoulders of the tension bars. In the other steels that showed cracks the cracks were about the same size as in 1158, but there were not as many, generally one or two. XI

(b) This shows the dendrites in Steel 1215, H. T., at higher magnification. X50



 α



6

Fig. 13.—Influence of considerable amounts of ferrite on tensile properties of heat-treated specimens, ×50

(a) Steel 1144. C, 0.38 per cent; Si, 1.35 per cent; Mn, 0.84 per cent; Ni, 3.10 per cent; Al, 0.005 per cent; Ti, 0.017 per cent; Zr, 0.32 per cent. This steel gave 307 000 lb./in.² tensile strength with 7.5 per cent elongation in 2 inches, and 21.7 per cent, reduction in area in the heat-treated bar. The structure is fine martensite

martensite
(b) Steel 1197. C, 0.32 per cent; Si, 1.37 per cent; Mn, 0.60 per cent; Ni, 3.05 per cent; Al, 0.02 per cent; Ti, 0.17 per cent; Zr, 0.32 per cent. This steel is of almost identical chemical composition as one shown in (a), but gave only 217 000 lb./in.² tensile strength with 3 per cent elongation in 2 inches and 27.5 per cent reduction in area. The carbon is slightly lower and may partly account for the large amount offerrite shown in the micrograph, but it is more probable that this is, because the specimen was not heated high enough before quenching

air-cooled specimens (see Fig. 8, e, f), but the steels also show characteristic inclusions. At a magnification of 500 diameters the two cerium steels that were examined were found to contain a large number of gray inclusions, and also some large circular orange inclusions, with interior light-gray markings. The uranium steels contained deep-blue inclusions (see Fig. 9). It seems that cerium and uranium, in a way somewhat similar to manganese (see Fig. 8, g), act both as true alloying elements and to produce soundness in the metal. Copper goes into solution, but did not produce a martensitic pattern in the air-cooled specimens, except in the one of high copper content, 1.35 per cent (see Fig. 8, h).

Boron forms a complex eutectic, probably that of an ironcarbon-boron compound with iron (see Fig. 10). The difficulty experienced in rolling the ingots containing boron is due to the fusibility of this eutectic at the temperatures ordinarily used for rolling. In the ingots that broke in the rolls the cracks in the polished specimens were found to follow the eutectic structure observed in the steels as cast, but in other places no traces were visible. Instead hard spherical particles, evidently of a single constituent of the same appearance as iron carbide, were found (see Fig. 11, a, b). The mechanical working probably breaks up the eutectic, the iron of the eutectic is absorbed by the iron of the matrix, while the iron-boron-carbon compound coalesces into the hard circular particles. These particles no longer form a weak brittle network and may have an appreciable hardening effect on the properties of the steel which may be desirable for the purpose of armor plate. Care must be taken, however, in the final heat treatment that the steel is not heated to too high a temperature before quenching, as the eutectic will then again appear and render the steel unsuitable (see Fig. 11, d). In the unetched state both the eutectic and the hard circular particles are pink, they both etch dark with sodium picrate, similar to simple iron carbide, and are not etched by 2 per cent nitric-acid etching, but appear white in contrast to the etched matrix (compare Fig. 11, a, b with Fig.

All of the above elements are true alloying additions; they are metallic constituents of the finished steel. The properties they confer upon the steel can not be established from a microscopic examination alone, but must be determined by physical tests. In the light of the considerations indicated above, however, they may all give good results, but more care perhaps must be taken

in the making and treatment of the cerium, uranium, and boron steels than of the others.

(c) SOUNDNESS OF THE STEELS AND STRUCTURES OF THE NORMALIZED AND HEAT-TREATED SPECIMENS

Not all the steels that were made up were examined under the microscope. Typical specimens, however, were chosen from every class of chemical composition, and all those specimens that gave exceptional or unexpected values in the tension tests were examined. Some samples were examined from the ingot as cast and some from the plates as rolled, but the great majority of the specimens were taken from the tested tension bars.

In most of the steels that contained zirconium and titanium in appreciable amounts there were found in the polished and etched specimens the thin, yellow streaks which are the segregates of the zirconium and titanium inclusions rolled out into plates (see Fig. 12, a, steel 1213). Many of the steels also showed small longitudinal cracks in the specimens (see Fig. 12, a, steel 1158), while in others there appeared a pronounced dendritic pattern (see Fig. 12, a, steel 1215, and Fig. 12, b). These features are of sufficient size as to be seen with the naked eye. The streaks and cracks, since they occur longitudinally in the specimens, should not be expected to have much effect in lowering the tensile properties as measured. The dendritic pattern, which of itself is considered undesirable, was not found to be associated particularly with steels that gave low values in tension. In fact, some of the steels which had the highest tensile properties showed a dendritic pattern. This is perhaps because these steels were rolled with the least number of reheatings, giving a better plate by avoiding repeated heatings and oxidation. The steels that were rolled in the latter part of the investigation, presumably most expertly, gave the largest percentage of dendritic structures.

The following is a list of these steels which showed small cracks in the specimens examined (see Fig. 12) and those which had a dendritic structure. The normalized steels are marked N. and the heat-treated ones H. T.:

```
      Cracked:
      Dendritic:

      1106, N.
      1103, N. and H. T.

      1109, N.
      1132, N. and H. T.

      1119, N.
      1136, N.

      1157, H. T.
      1167, N.

      1158, N. and H. T.
      1168, N. and H. T.
```

Cracked—Continued	Dendritic—Continued
1176, H. T.	1190, N.
1178, N.	1200, N.
1189, H. T	1205, N.
1211, H. T.	1206, N.
1213, N.	1207, N. and H. T.
1228, N. and H.·T.	1215, N. and H. T.
1229, H. T.	1216, N.
1236, N. and H. T.	1219, N. and H. T.
1237, H. T.	1224, N.
1258, N. and H. T.	1227, N. and H. T.
1274, N. and H. T.	1231, N.
1275, N.	1237, N.
1278, N. and H. T.	1244, N. and H. T.
1286, N.	1252, N.—Streaky.
	1285, N.
26-	1286, N. and H. T.

The normalized, as well as the heat-treated specimens, were partially decarburized uniformly to the depth of 0.005 inch along the gage length of the tension specimen, which makes 0.010 inch of the diameter of the gage length decarburized. The differences in the microstructures of the air-cooled specimens have already been described. In the heat-treated specimens those that gave the best results in the tension tests invariably had a structure of fine martensite. Wherever appreciable amounts of ferrite were found, together with the martensite in the quenched and tempered specimens, the tension values were not so good. In many cases unexpected low results could be traced to the presence of relatively large amounts of ferrite (see Fig. 13). The variations in the drawing temperatures, if there were any, could not be detected in the microstructure.

Altogether, in this part of the examination 160 micrographs of the steels in the cast, rolled, and finally heat-treated conditions were taken. For lack of space they can not be given here, but they were invaluable in helping to interpret the results of the mechanical tests. By means of this survey of the microstructures of the samples those containing flaws were discovered and eliminated, and any irregular results could be explained. The microscopic "check-up" thus served to relieve any doubts concerning single samples and helped to give weight to the general conclusions of the investigation.

3. MECHANICAL PROPERTIES

(a) TENSILE TESTS

Since the rolled material was one-half inch or less in thickness, it was impossible to use the standard form of tensile specimen having a diameter in the reduced section of 0.505 inch. A shoulder type specimen was used, having a diameter of 0.300 inch at the reduced section and a gage length of 2 inches, as shown in Fig. 14. This ratio of gage length to area of specimen gives values of elongation somewhat less than would be obtained with standard size specimens, and this fact should be borne in mind in interpreting the results. The tensile and also the impact specimens

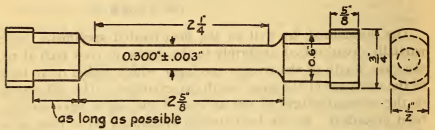


FIG. 14.—Dimensions of tensile specimens

were completely machined before heat treatment and not ground before testing.

Tensile tests were made on either a 50 000-pound or 100 000-pound Riehle testing machine. Proportional limit was determined from plotted stress strain curves, strain being measured by means of a Berry strain gage. In a few cases the specimens were too short to admit of fastening a strain gage to the specimen, and blanks appear in the tables (Tables 11-22) for these cases. Yield point was determined by the "drop-of-the-beam" method where any "drop" was observed. Reduction of area and elongation were obtained by the usual measurements after fracture.

Tables 11-22 give the results of the tensile tests on all the steels. It will be noted here that several of the steels have a tensile strength of well above 300 000 lbs./in.², accompanied by appreciable ductility. No. 1207, with a tensile strength of 344 000 lbs./in.², was the highest value observed.

TABLE 5.—Hardness Measurements on Plate

[*Indicates specimen not completely broken]

	1		posimon i	lot completely broken]				
	This-t-	Hardness	numerals		Thirt	Hardness	umerals	
Ingot No.	Thick- ness	Brinell	Sclero-	Ingot No.	Thick- ness	Brinell	Sclero-	
		Billiell	scope			Brillen	scope	
	Inch				Inch			
1101	3/8 1/2	334 293 405 444	30 26 52 47 27 25	1159	3/8/2/8/2/8/2/8/2/8/2/8/2/8/2/8/2/8/2/8/	294 327 603 477	32 32 55 45	
1102	3/8 1/2	240 245 217 223	21 22	1160	3/8 1/2	573 279 556 545	60 52	
1103	3/8/2/8/2/8/2/8/2/8/2/8/2/8/2/8/2/8/2/8/	207 207 223 212	32 31	1161	3/8 1/2	285 440 432 512	24 42 31 47	
1104	3/8 1/2	207 194 187 205	20 21 20 22	1162	% 1/2	321 337 387 338	24 23 38 37	
1105	1/2	172 241 232 207	28 28 30 28	1163		219 234	21 22	
1106	3/8	189 235 202 216	24 25 21 26	1164 1165*		375 340 338 346	30 28 36 40	
1107	3/8 1/8	189 182 187 196	22 20 20 21	1165* 1166. 1167. 1168.		430 514 364 387	33 32	
1109	3/1/3/1/3/1/3/1/3/1/3/1/3/1/3/1/3/1/3/1	236 231 244 255	24		i	1	44 43	
1111	3/8 1/9	317 335 340 351	30	1169. 1170. 1171. 1172. 1173.		640 571 550 600	61 57	
1112	3/8	415 302 290 438	41 31 39 40	1171		387 447 424 428	34 41 43 37	
1113	_	467 244	39 34				28 26	
1114	3/1/3/1/2/3/1/3/1/3/1/3/1/3/1/3/1/3/1/3/	477 484 537 491	43 43 48 49	1174 1175* 1176.		226 206 472 467	19 19 52 52	
1115	1/2 3/8	364 354	62 60 34 37	1176 1177* 1180		470 481 477 477	36 38 55 55	
1117	3/8 3/8	361 340 626 508	47 42 43 43 57 68		1		19 23	
1118	3/8	600 576 626 626	57 68	1181		208 216 213 224	20 21 20 21	
1110	1 -	387 364	42 40	1182. 1183. 1184. 1185.		257 241 171 172	18 19	
1119	3/1/3/1/3/1/3/1/3/1/3/1/3/1/3/1/3/1/3/1	555 600 495 470	54 52	1186				
1128	1/2 3/2	418 495 380 370	52 43 47 52 35 36	1187. 1188. 1189*		381 403 300 311 342 336		
1129	1/2 3/6	467 418 444 387	42 45 39 42	1189*. 1190.		351 360 351 325	42 44 34 29	
1130	1/2	475 460 387 370	47 42 37 35 37 35	1191		344 338	1	
		338 375	37 35	1192		228 217	24 23 26 26	
1131	8/8 1/2	652 635 575 534 558 552	53 63	1193. 1194. 1195.		439 457 444 403	52 53 42 38	
1132	8/1/8/21/8/21/8/21/8/21/8/21/8/21/8/21/	589 626	55 65				46 34	
1133	3/8	387 488 444 402	40 54 45 40	1196. 1197. 1198.		415 369 209 218	36 36 23 24	
1134	3/8 1/2	477 470 444 477	42 48 55 58	1198. 1199. 1200.		216 210 321 302	24 23 37 28	
1135	3/8 1/2	444 336 512 444	47 43	1201		268 266	26 26	
1136	3/8	491 479		1201 1202 1204*		207 199 477 457	21 21 50 53	
1138	3/8	410 375 589 608	37 34	1205. 1206.		505 512 481 491	41 45 42 47	
1144	3/8	578 591 578 346 589 555	50 56 53 26 55 50	1207		627 600 629 616	55 55 69 61	
1145	3/9	351 245 444 352	35 25	1208. 1209. 1210. 1211.		629 616 600 555 378 411	66 62	
1146	90/01/00/00	418 390 478 321	41 31 32 32 45 28	1211		456 447	46 40	
1147		487 450	52 60	1212* 1213		367 361 471 564	43 45 42 51	
1155	8/1/8/1/8/1/8/1/8/1/8/1/8/1/8/1/8/1/8/1	387 477 585 555	37 45	1214 1215 1216		481 495 600 553	41 37	
1156	1/2	519 550 477 516	51 53 43 45			418 387	34 35	
1157	1/2 3/8	537 534 418 423	56 55 38 39	1217. 1218.		475 477 310 278	55 55 30 28	
1158	1/2 8/8	518 522 544 640	55 59	1219		235 228 441 337	23 23 45 36	
	1/2	452 452	50 51	1221		312 323	33 33	

TABLE 5.—Hardness Measurements on Plate—Continued

[*Indicates specimen not completely broken]

	(T)	Hard	ness	nume	erals			Hardı	1ess r	ume	rals
Ingot No.	t No. Thickness	Bris	nell				Thick- ness			Sclero- scope	
222*	Inch	366	367	47	49	1259	Inch	596	605	47	48
223		357	317	33	26	1260		532	569	57	48
224		406	438	32	34	1261		207	223	18	19
225		467	387	32	41	1263		402	373	37	41
226		430	454	36	37	1264	• • • • • • •	387	396	43	4
227		550	537	54	54	1267*		180	183	27	2
228		629	627	66	62	1268		252	219	23	2
229		585	573	57	55	1269*		207	188		
230		603	605	70	69	1270*		351	364	20	2
231		520	474	47	42	1271*		341	342	38	4
232		494	505	44	46	1272*		250	321	25	2
233		321	311	28	26	1273*	• • • • • • •	255	216	32	2
234		364	375	30	30	1274*		444	418	52	4
235		351	241	23	22	1275*		555	600	55	6
236		484	472	54	53	1276*		534	570	52	5
	1										
237		600	585	56	54	1277*		410	444	42	5
238		444	474 555	46	49	1278*		395	397	44	4
239 240		532 509	516	59 41	61 42	1279*. 1280*.		490 510	520 486	50	6
241		520	512	47	47	1281*		508	468	60	5
		320	312	47	17	1201		300	700	00	3
242		387	430	39	41	1282*		510	475	57	5
243		460	438	42	43	1283*		539	512	57	5
244		441	420	44	45	1285*		475	430	49	4
245*		525	493	56	57	1286*		520	508	54	5
246		444	444	39	36	1289*		240	448	-00	4.
247		366	418	33	36	1290*		340 455	447 498	38 51	5
248		248	256	20	20	1291*		464	478	50	4
249		361	371	19	18	1292*		522	486	60	6
250		324	324	28	26			418	470	39	4
251		477	495	39	37						
252		477	487	40	39						
253		600	560	53	49						
256		532	495	49	49						
257		524	477	52	47						
258		441	440	38	40						

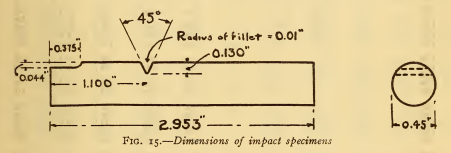
(b) IMPACT TESTS

Impact tests were made on heat-treated specimens from all heats beyond No. 1155 and a few previous, the heat treatment being the same as for the tensile specimens and plates. Hardness tests made on a few impact specimens gave sufficient evidence that the impact specimens were in a structural condition similar to the tensile specimens. An Izod machine using a cantilever type of specimen was used in all impact tests.

Here, again, the thickness of the available material precluded the use of the standard type of specimen in all cases. For those plates which would not admit of making a round specimen 0.450-inch diameter (see Fig. 15), the largest diameter possible in multiples of 0.050 inch, was used and the notch made geometrically similar to the larger specimens. The total length of the specimens re-

mained constant, so as not to alter the striking distance. The specimens of small size were held in the anvil of the impact machine by means of hardened steel split sleeves having an outside diameter of 0.450 inch and an inside diameter to fit the specimen. This sleeve was inserted in the anvil flush with its top surface and the impact specimens properly aligned by means of a templet. The height of fall of the pendulum was varied proportionally to the size of the specimen, although theoretically this should not be necessary.

The area of the specimens at the notch was computed and the energy absorbed in breaking for unit area determined, a value which has been called the specific impact work. The results of the tests are given in Table 6. While the law of similarity has not been definitely shown to hold for impact specimens as for other



forms of mechanical testing, the method used was considered the most desirable that the circumstances would permit.

A high impact value does not always indicate a superior steel, since such values are many times accompanied by low tensile strength. Thus, all the steels which had an impact value greater than 200 foot-pounds per square inch also gave tensile strengths less than about 130 000 lbs./in.², except No. 1252, which had a tensile strength of 324 800 lbs./in.². On the other hand, if we select all those steels which showed an impact value less than 50 we note that the majority of that group have a tensile strength in excess of 275 000 lbs./in.², although a few fall even below 100 000 lbs./in.², the latter being clearly inferior steels. Those steels, then, that show fair values of impact, together with high tensile strength, should be considered the best steels, since they combine strength with toughness.

TABLE 6.—Impact Tests (Izod Machine)

[*Indicates specimen not completely broken]

			7-7	-	
	Diame-	A =00 =4	Initial	Energy a	hanrhad
Ingot No.	ter of specimen	Area at notch	energy	in bre	aking
	specimen				
- n - M					
	Inch	T	Ftlbs.	Ftlbs.	Ft Ibs./in. ²
1135	0. 450	Inch ² 0. 1210	120	7. 0	58
1136	. 450	. 1210	120	8.0	66
1138 1155	. 447	. 1196	120 120	6. 5 13. 5	54
1156	. 447	. 1196	120	5. 0	113 42
	1	.1150			
1157	. 450	. 1210	120	13. 5 5. 5	112
1158 1162	. 448	. 1201	120 120	6. 0	46 51
1163	. 448	. 1201	120	29. 5	246
1164	. 347	. 0732	75	8. 0	109
1165	. 449	. 1206	120	9. 5	79
1166	. 346	. 0730	75	8.0	110
1167	. 350	. 0736	75 120	4. 5	61 99
1169	. 450	. 1210	75	12. 0 5. 5	75
		1			
1170		. 0738	75	4.5	61
1171 1172	. 454	. 1229	120 75	16. 5 11. 0	134 148
1173		0741	75	9. 0	121
1174	. 449	. 1206	120	16.0	133
1175	. 447	. 1196	120	10.5	88
1176	. 354	. 0742	75	8.0	108
1177 1178			120	9.3	78
1180	. 450	. 1210	120	14.0	116
1181 1182		. 0524	60 120	6. 5 15. 0	124 126
1183	349	. 1187	75	6.0	82
1184	. 444	. 1182	120	28. 0	237
1185	. 446	. 1192	120	42. 0	357
1186	. 441	. 1168	120	19.0	163
1187	. 349	. 0735	75	9.0	122
1188 1189		. 0733	75 75	10. 0 8. 0	136 109
1190	. 450	. 1210	120	14.5	120
					101
1191 1192		. 0742	75 120	7. 5 24. 5	101 202
1193	. 354	. 0742	75	17. 5	236
1194	. 447	. 1196	120	10. 5	88
1195	. 349	. 0735	75	7. 5	102
1196	. 347	. 0732	75	9. 5	130
1197	. 348	. 0733	75	8. 5 75. 5	116 *625
1198	. 449	. 1206	120 75	33. 5	*460
1200	. 447	.1196	120	14.5	121
1201	. 349	0725	75	16.0	218
1202	. 452	.0735	120	41.5	340
1204	. 449	. 1206	120	12.0	100
1205 1206	. 350	. 0736	75 75	7. 0 9. 5	95 129
	. 349	.0/35	/3		
1207	. 350	. 0736	75	5. 5	75
1208 1209	446	. 1192	120 75	4. 5	38 20
1210	. 446	.1192	120	14. 0	117
1211	. 348	. 0733	75	9. 5	130
1212	. 447	. 1196	120	12.0	100
1213	. 447	.1196	120	4.5	38
1214	. 349	.0735	75	6. 5 6. 0	. 88 50
1215 1216	. 447	.1196	120 120	13. 5	112
1217	. 350	.0736	75 75	6. 5 9. 0	88 122
1219.	. 448	. 1201	120	17. 5	146
1220	. 447	. 1196	120	9.0	75
1221	. 353	. 0741	75	9.0	121

TABLE 6.—Impact Tests (Izod Machine)—Continued

[*Indicates specimen not completely broken]

Vanish No.	Diame-	Area at	Initial	Energy absorbed		
Ingot No.	ter of specimen	notch	energy	in breaking		
	-111				Ft	
	Inch	Inch ²	Ftlbs.	Ftlbs.	lbs./in.2	
1222. 1223.	0.450	0. 1210	120	11.0	91 79	
1223	. 449	. 1206	120	9. 5 7. 5 8. 5	101	
1225	. 349	. 0741	75 75	8. 5	116	
1226	. 298	. 0534	60	4.5	83	
1227	200	0520	60	4.0	74	
1228	. 299	. 0539	60	4. 0 2. 0	37	
1228. 1229.	. 348	. 0539	75	8. 0	109	
1230	. 296	. 0524	60	. 5 8. 0	10 109	
1231	. 350	. 0736	75	8. 0	109	
1232	. 299	. 0539	60	4. 0	74	
1233	. 300	. 0544	60	4.0	74 74	
1233	. 349	. 0735	75 75	6. 5 13. 5	88	
1236	. 350	. 0736	75 75	13. 5 8. 5	183 116	
2000	. 349	. 0735			110	
1237	. 351	. 0733	75	4. 5 4. 5	61	
1238	. 298	. 0534	60	4. 5	84	
1240	. 299	. 0539	60 60	1. 0 6. 5	19 120	
1238. 1239. 1240. 1241.	. 299	. 0534	60	1.5	28	
1242 1243	. 351	. 0738	75 60	6. 5	88 138	
1244	. 300	0735	75	7.5	138	
1245 1246	. 299	. 0735	75 60	7. 5 7. 5 3. 5	65	
1246	. 300	. 0544	60	2. 5	46	
1247	. 350	0726	75	4. 0	54	
1247	. 351	. 0736 . 0738	75	6. 0	81	
1249	349	. 0735	75 75	10.0	136	
1250	. 299	. 0735	60	5. 5 2. 0	102	
1451	. 299	. 0539	60	2. 0	37	
1252	. 298	. 0534	60	13. 5	252	
1253	. 298	. 0534	60	2. 5 21. 5	47 174	
1257	. 450	. 1210	120 100	21. 5	174	
1258	. 350	. 0736	75	6. 0 5. 0	72 68	
		1				
1259	. 399	. 0827	100	8. 0	97 72	
1260	. 400	. 0833	100 120	6. 0 5. 8	48	
1261	. 400	. 0833	100	4. 0	48	
1264	. 450	. 1210	120	5. 5	46	
1267	. 398	. 0821	100	3. 0	37	
1268	- 400	. 0833	100	6. 0 14. 5	72	
1269	. 449	. 1206	120	14. 5	120	
1270	. 398	. 0821	100 120	10. 5 10. 5	128 91	
12/1		. 1200	120	10. 3	91	
1272	. 397	. 0815	100	10. 5	128	
1273. 1274.	. 397	. 0815	100	31.0	380	
1275	. 389	. 0827	100 120	3. 0 5. 0	36 41	
1275	. 450	. 1210	120	4. 2	35	
1277		0000	20.00			
1277	. 348	. 0733	75 120	8. 0 19. 5	109	
1278. 1279. 1280.	. 448	. 1210	120	4.0	34	
1280	. 448	. 1201	120	6.0	161 34 50	
1281	. 449	. 1206	120	4. 0	33	
1282	. 450	. 1210	120	7. 0	58	
1283	. 449	. 1206	120	3.0	58 25 83	
1286.	. 449	. 1206	120	10. 0	83	
1283. 1285. 1286. 1289.	. 349	. 1206	120 75	8. 0 7. 0	66 95	
	1 1 1					
1290	. 350	. 0736	75	6.0	82	
1292	. 349	. 0735	75 75 75	5.5	75 75 95	
1293	. 349	. 0735	75	5. 5 5. 5 7. 0	95	
	1					

(c) HARDNESS TESTS

Hardness tests were made on the tensile specimens and on the hardened plates. For the tensile specimens this consisted of two Brinell impressions on the ends of the broken specimens after grinding off the outside surface. Scleroscope hardness was also determined on the same material, several readings being taken with a recording scleroscope.

On the plates opposite corners were ground down and duplicate Brinell and several scleroscope determinations made at each posi-

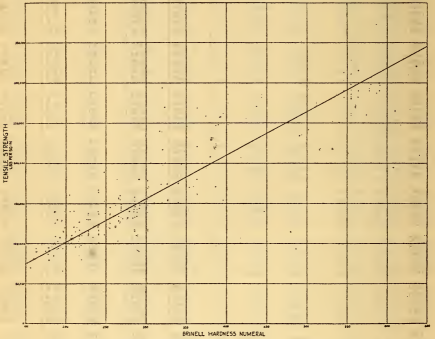


Fig. 16.—Relation between tensile strength and Brinell hardness

tion. In this case an indicating scleroscope was used for some of the work and a recording instrument for the remainder. The indicating instrument gave uniformly lower values than the recording type, so that the scleroscope values for the plates are not in all cases intercomparable. Those taken with the latter instrument are marked with an asterisk in Table 5, which gives the hardness values for both corners of each plate.

The method of heat treatment described in Section III-2 does not give exactly the same hardness to both the small tensile specimens and the relatively larger plates. To secure the same hardness, it would have been necessary to draw back the tensile specimens at varying temperatures until like conditions were reached.

Burgess Woodward]

By aid of

By aid of Fig. 16, showing the relation between tensile strength and hardness for this class of steels, an idea of the actual tensile properties of the hardened plates may be obtained.

V. COMPARISON TESTS ON SIMILAR MATERIAL

In addition to the ingots prepared by the Bureau of Mines there was also available for study other material of a similar nature which was submitted through the Bureau of Ordnance of the Navy Department and was secured from one of the large automobile manufacturers who was constructing armored tanks during the war and whose representatives had made great claims for zirconium as an alloying element in light armor plate. This material consisted of 45 plates of ½ to ½ inch thickness, representing 28 separate heats of steel. Each heat, comprising about 1000 pounds of metal, was made in an electric furnace.

The majority of the plates as submitted were 18 inches square. The following, however, were 12 inches square: 16–1, 19–1, 20–3, 22–2, 22–3, 22–4, 24–5, 25–1, 25–2, 25–4, 25–8, 27–3, and 27–4. (The first number in all cases refer to the heat number and the second to the plate number of that heat.) The plates were supposedly heat treated, but many were found to be soft and were heat treated at the Bureau of Standards in accordance with a summary of the heat treatments given the same material by the manufacturer.

All plates were cut up, the hardened ones by grinding, so as to produce two tensile bars, an impact specimen, and a 12 by 12 inch ballistic plate from the large size plates. From the smaller plates a ballistic plate of 11 by 11 inches was obtained in most cases.

In Table 7 is given the heat treatment of the plates and test pieces on those plates which were heat treated at the Bureau of Standards.

The same type of specimen and testing procedure was used for these materials as for those described above, with the exception that the B and C specimens are both hardened and no tests were made on normalized material.

In Table 12 will be found the results of impact tests calculated in a similar manner to those given before.

A comparison of these tests made at the Bureau and those made by the manufacturer is given in Table 9, the average value for each heat being given. In the grand average certain of the heats are omitted, as noted in the table, since the figures from both sources are not strictly comparable.

TABLE 7.—Comparison Steels

[AC=Air cooled; FC=furnace cooled. All specimens drawn for one hour in oil bath unless otherwise stated]

TT 4	Te	mperature	s			Te	mperature	S	
Heat No.	Normal- izing	Quench- ing	Draw- ing	Remarks	Heat No.	Normal- izing	Quench- ing	Draw- ing	Remarks
2 4 6 8 9 10 11 12 13 14	870 AC 870 AC 870 FC 870 AC	850 850 850 850 850 830 777 843 850 843	° C 205 205 205 193 193 177 205 228 290 193	Salt bath for drawing.	18 19 20 21 22 23 24 25 27 28	870 FC 816 AC 954 AC 870 FC 870 FC 870 AC 870 AC 900 FC 800 AC 816 AC 800 AC	850 860 850 850 850 850 850 850 850 800 843 800	° C 190 190 316 190 205 205 200 200 200	Salt bath for drawing Drawn for three hours

a These values not given by manufacturer, but estimated at Bureau of Standards.

Table 10 gives the results of hardness measurements on ground portions of opposite corners of the plates. The results that were obtained on these comparison steels are quite similar to those obtained on our own material, but it will be noted that the highest tensile properties observed were not so great as those from the regular series. The comparison steels will therefore be included in the discussion of results and the effect of the various elements on the properties of steel.

TABLE 8.—Impact Tests (Izod Machine) of Comparison Steels

Steel	Diameter of spec.	Area at notch	Initial energy	Energy a	
	4 11	10.0			Ft
	Inch	Inch 2	Ftlbs.	Ftlbs.	lbs./in.2
2–1	0.449	0. 1206	120	17. 0	141
4-1	. 397	. 0815	100	6.0	74
6-1	. 448	. 1201	120	4.0	34
8–1 ,	. 399	. 0827	100	12. 0	145
9–1	. 449	. 1206	120	4.0	33
9-2	. 400	. 0833	100	3, 5	42
10-2	. 450	. 1210	120	14. 0	116
11-2	, 400	. 0833	100	14. 0	168
12-1	. 399	. 0827	100	13. 0	157
13-2	. 499	. 1206	120	6.0	50
	399	. 0827	100	15.0	101
14-1			100 100	6.0	181 72
15-1	. 400	. 0833	75	3. 0	41
16-1	. 450	. 1210	120	15. 0	124
18-1	. 349	. 0735	75	7. 0	95
19–1			,-		
20–1	. 348	. 0733	75	5. 5	75
20–3	. 449	. 1206	120	6.0	50
22–1	. 349	. 0735	75	8. 0	109
22-2	. 450	. 1210	120	16. 5	136
22–3	. 449	. 1206	120	9. 0	75
22-4	. 449	.1206	120	9.0	75
23-1	. 349	. 0735	75	9.0	123
24–5	. 348	. 0733	75	7.0	96
25–1	.398	.0821	100	11.0	134
25-2	. 449	. 1206	120	9.0	75
25–4	. 398	. 0821	100	10, 5	128
26-1	. 349	. 0735	75	2. 0	27
27–3	. 301	. 0549	60	4. 0	73
27-4	. 349	. 0735	75	8. 0	109
28-1	. 450	. 1210	120	4. 0	33
	3 100	1210			

TABLE 9.—Comparison Steels

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			Bureau	Bureau of Standards' tests	s' tests			Mar	Manufacturers' tests	tests	
Kind of steel	Heat Nc.	Tensile	Yield point	Elongation in 2 inches	Reduction in area	Brinell hardness numeral	Tensile	Elastic limit	Elongation in 2 inches	Reduction of area	Brineli hardness numerai
NI, SI, Zr. NI, SI, Zr. Cr, V, Mo. NI, SI, Co, Mo. NI, SI, Cr, Zr.	∺ 0.004.0	Lbs./in.² 257 100 228 800 288 600 253 200 308 800	Lbs./in.² 188 800 202 900 229 500 238 500 247 900	Per cent 11.3 9.6 7.8 7.8 8.8	Per cent 37.3 35.3 35.2 26.2 2 20.1	523 436 589 556 615	Lbs./in. ² 301 000 272 000 304 000 202 000 285 000	Lbs./in.² 250 000 231 000 264 500 202 000 263 000	Per cent 10.5 10.5 11.8 11.0 2.0	Per cent 25.9 31.5 33.8 33.8 3.3 6.4	555 477 567 532 535
Or, V, Mo. Ni, Si, Zr. Si, Zr. Or, V, Mo. Ni, Si, Zr.	a 6 6 7 7 8 8 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	237 200 282. 500 245 000 186 500 285 600	209 900 254 700 202 800 259 400	, 4, 5, 5, 5, 5, 5, 5, 5, 5, 5, 5, 5, 5, 5,	29.0 29.0 31.6	540 584 440 556	(b) 294 000 265 000 183 000 225 000	(b) 264 000 210 000 200 000	6. 5 16. 5 1. 0 10. 0	8.8 16.5 1.0 36.0	600 555 512 512 566 532
NI, Mo, Zr NI, Si, Cr, Zr NI, Co, Si, Zr NI, Co, Si, Zr NI, Si, Co, Mo	111 12 13 14 14	208 700 283 300 172 700 325 400 226 700	187 800 221 600 (c) 268 300 (c)	7.8 (°) (°)	28.6 33.3 19.8	324 445 545 534 509	202 500 295 500 284 000 281 000 260 000	176 500 255 000 250 000 238 000	13.5 4.5 10.0 10.3 1.0	46.5 9.0 30.0 32.0 5.0	555 578 545 578 578
N, S, Z N, S, Z N, S, Z N, C, S, Z N, S, Z N, S, Z	a 16 a 17 118 119 20	236 800 (d) 253 900 262 600 301 800	226 200 (d) 208 800 227 800 236 900	(d) 11. 5 11. 5 5. 7	(d) 32.9 33.3 14.9	564 555 567 542	(b) 272 500 285 700 287 000 312 000	(b) 233 500 250 000 254 500	8.5 10.7 3.0 10.3	27. 0 30. 7 5. 0 30. 0	512 477 543 555 534
Ni, Si, Zr. Ni, Co, Si, Zr. Ni, Si, V. Ni, Si, Zr. Ni, Si, Zr.	21 22 23 23 24 24	252 500 277 900 292 300 281 500 303 600	245 800 238 100 224 600 228 800 245 100	10.0 8.7 8.7 9.0	28. 33.3.4.4 33.3.4.4	424 512 512 578 600	270 000 290 000 326 000 297 000 309 000	235 000 275 000 277 000 242 000 245 000	12.5 11.0 7.8 8.5	44. 0 43. 0 16. 0 21. 5 29. 0	512 512 534 532 544
Ni, Si, Zr. Ni, Si, Zr. Ni, Si	a 26 27 a 28	(e) 283 800 (e)	(c) 254 200 (c)	7.9	31.9	636 385 593	(e) 295 000 (e)	(e) 250 000 (e)	2.5	4. 5	009
Average		269 700	230 600	7.5	23. 5	514	275 300	239 800	8.3	22. 6	536
a Values for these heats not included in average.	l in average.	b Broke	b Broke in head.	c Broke	e Broke in shoulder.	d Not t	d Not tested in heat-treated condition.	-treated con	idition.	e No values given	given

TABLE 10.—Comparison Steels [Hardness of plates on opposite corners]

	Har	dness	num	eral		Hard	iness	num	eral
Plate No.	Brit	ıell		ero- ope	Plate No.	Brit	ielí	Scle	ero-
1-32-1	532 418	495 402	61 49	a 60 47	17-1 18-1	262 512	262 526	29 60	a 36
2-2. 3-1. 4-1	375 607 510	387 532 495	62 53	a 45 a 61 51	18-2	273 486	277 522	32 58	a 5
4-2					19-2	293	293	32	a 33
5–1	311 600 600	311 546	38 57	a 38 a 55	20-1	504 514	516 571	56 57	6
5-2 6-1	555 477	600 578 495	65 63 55	a 65 67 a 57	21-1	- 474 514	470	58	5 5
6–2 7–1	600	555		a 55	22-1 22-2 22-3	553 495	479 512	61 67 58	5 4
7-28-1	512 474	495 452	58 62 53	a 60 52	22-4	474	474 465	47	4
9–1 9–2	560 555	555 568	66 61	65 63	23–1 24–5	571 532	544 483	72 64	6
0-2	508	512	57	63	25-1	522 529	518 553	56 59	5
1–2	418 509	418 512	56 65	53 67	25–4	510	526	57	6
3–1	258 567	262 522	33 66	a 35	25-8 26-1	544 562	550 562	65 71	6
4–1	504	500	66	59	27-3. 27-4.	407 439	446 452	47 53	4 5
5–1 6–1	512 477	486 460	60 50	59 49	28–1	522	480	-60	5

a Plates not heat treated at Bureau.

VI. EFFECT OF VARIOUS ADDITION ELEMENTS

The large number of steels examined offers an excellent opportunity for studying the effects of the various alloying elements. In nearly all the heats, however, the silicon content is higher than that usually obtained in ordinary practice. For purposes of comparison it is necessary to classify the steels into groups having the least possible number of variables in each group. The carbon content is a variable in practically every group, and the arrangement given in the tables is according to increasing carbon content.

The groups into which the steels have been roughly classified are as follows: Group A, silicon steels; Group B, nickel-silicon steels; Group C, silicon-zirconium steels; Group D, nickel-silicon-zirconium steels; Group E, cerium steels; Group F, copper steels; Group G, boron steels; Group H, uranium steels; Group I, molybdenum steels; Group J, nickel-chromium steels; Group K, vanadium steels; Group L, chromium-tungsten steels; Group M, cobalt steels.

1. GROUP A-SILICON STEELS

This group, shown in Table 11, represents plain carbon steels in which the silicon is greater than normal and which have all been deoxidized with aluminum. The group has been further divided into steels that have greater or less than 1 per cent silicon. It will be noted that the increase of silicon to above 1 per cent has resulted in a greater tensile strength and impact value without materially reducing the ductility. Nos. 1269 and 1270, containing, respectively, 0.65 per cent titanium and 0.45 per cent aluminum, show no superiority over No. 1104, which is simply deoxidized with these elements.

2. GROUP B-NICKEL-SILICON STEELS

Table 12 illustrates this group which has been further classified into steels containing 2 per cent nickel, 3 to 3.25 per cent nickel with silicon greater or less than 1 per cent, and those having more than 3.25 per cent nickel.

The class with 2 per cent nickel all contain approximately 1 per cent silicon and show increased mechanical properties in comparison with the corresponding class of Group A. A few steels in this class have tensile strength in the neighborhood of 300 000 lbs./in.², but the ductility and toughness are not as great as in those that follow.

The 3 per cent nickel group again shows the advantage of increasing the silicon to greater than 1 per cent. In fact, this combination of elements represents about the best of any of those tested, the majority having a tensile strength from 270 000 to 315 000 lbs./in.², depending upon the carbon content, yield point from 200 000 to 250 000 lbs./in.², and proportional limit from 100 000 to 160 000 lbs./in.², excellent ductility and satisfactory impact values. The values for the normalized steels are also excellent. A carbon content of from 0.40 to 0.50 seems to be the most favorable.

The nickel content apparently should be kept in the range 3 to 3.25 per cent, as those steels having a higher percentage than this were nearly all brittle.

Group B also shows, as did Group A, that aluminum and titanium in amounts greater than that needed for deoxidation offer no advantage and, in fact, appear to be in most cases detrimental.

3. GROUP C-SILICON-ZIRCONIUM STEELS

This group as tabulated in Table 13 should be compared with Group A, which contains similar steels without zirconium. The zirconium content is variable from a small amount to 0.60 per cent. A study of Tables 11 and 13 seems to indicate that in the steels of lower carbon content the zirconium may have increased the ductility but not the tensile strength for the heat-treated steels. In the higher carbon range the ductility is much less than for similar steels in which zirconium is absent. The normalized steels containing zirconium have lower proportional limit, yield point, tensile strength, and ductility than those of Group A.

4. GROUP D-NICKEL-SILICON-ZIRCONIUM STEELS

These steels shown in Table 14 are subclassified similar to those of Group B. The 2 per cent nickel steels do not give as great tensile strength with zirconium as without it, but seem to show greater ductility and toughness. The same may be said of the 3 per cent nickel steels, although in this case the ductility is not increased to as great an extent, if any. There are several places in the table where the carbon content is constant and zirconium practically the only variable. None of these instances, however, show any regular effect on the properties attributable to the zirconium content.

5. GROUP E-CERIUM STEELS

The cerium steels without nickel, as shown in Table 15 and compared with the first portion of Group H, indicate that with about 0.25 per cent of cerium the tensile properties are increased with accompanying loss of ductility. The nickel-cerium steels have been arranged partly in order of cerium content in the table, since they are all of approximately the same carbon content. Although nearly all of the nickel-cerium steels have the nickel-silicon ratio shown in Group B to be desirable, it also appears that small amounts (up to 0.10 per cent) of cerium are beneficial, while larger quantities offer no further advantage. No. 1260, with only 0.01 per cent cerium, is a most excellent steel.

The rôle of cerium is thought to be that of a desulphurizer, and in Nos. 1256 and 1257 the sulphur content was intentionally increased to investigate this point. The tensile strength of these two steels is quite high, but the ductility is less than would be expected from a consideration of the other constituents. In amounts over 0.30 per cent cerium segregates very badly, and accordingly it would appear preferable to keep it below this figure.

6. GROUP F-COPPER STEELS

The copper has been added to these steels in place of a portion of the nickel content, and an inspection of Table 16 indicates that in those steels in which the sum of the nickel and copper, together with the silicon and carbon, are in the favorable ratio the usual high tensile strengths are secured, but with a reduction of the ductility and toughness. Specimens Nos. 1279 and 1280, for instance, broke in the shoulders, while the impact values are mostly low. No. 1285, with 0.70 per cent zirconium, apparently corrected for some of the lost ductility.

7. GROUP G-BORON STEELS

This group can almost be dismissed from further consideration because of manufacturing difficulties in producing sound steel containing boron. The majority of the steels (see Table 17) were of low carbon content, but do not compare favorably with similar steels of Groups H and B. The ductility is in all cases low even with small amounts (0.02 per cent) of boron. No. 1276, containing the favorable ratio of carbon, silicon, and nickel, did not show the high properties for that class.

8. GROUP H-URANIUM STEELS

There were only three steels containing uranium—Nos. 1228, 1229, and 1244—all having the favorable ratio of carbon, silicon, and nickel except No. 1228, which carried 0.63 per cent carbon. No 1244 showed the usual high properties, but the other two were less desirable (see Table 22).

9. GROUP I-MOLYBDENUM STEELS

With the possible exception of Nos. 3, the molybdenum steels (see Table 18) do not show the remarkable ductility claimed elsewhere for this element.⁵

This may, of course, be due to type of heat treatment to which all of these steels were subject, but it is probable that all of the steels in the nickel-molybdenum series would have been superior for the purpose desired with the molybdenum omitted.

⁵ Molybdenum as an alloying element in structural steels, G. W. Sargent, Proc. Am. Soc. for Test. Mats. 20, Part II, p. 5; 1920.

^{63593°--22---3}

10. GROUP J-NICKEL-CHROMIUM STEELS

The nickel-chromium steels classified in Table 19 all show good properties particularly regarding ductility and toughness. Although most of the steels contain zirconium also, the previous considerations would not indicate that this element has greatly influenced the results. The properties of all of the steels in Group J could be reproduced without the addition of either chromium or zirconium.

11. GROUP K-VANADIUM STEELS

This group contains the steel (No. 1207) which showed the highest tensile strength observed in the entire investigation—about 344 000 lbs./in.². Although the ductility is not so great as in certain of the other steels having very high tensile strength, it is nevertheless considerable for such a steel. In passing it might be worthy of mention that from a portion of the plate from this heat was constructed a spring for a precision aeronautic altimeter. This spring is constantly operating under a computed maximum fiber stress of 100 000 lbs./in.² and shows no elastic hysteresis or aftereffect, which is common to springs in such instruments. The group, as a whole, shows good properties but can not be considered as preferable to Group B (see Table 20).

12. GROUP L-CHROMIUM-TUNGSTEN STEELS

This group, consisting only of Nos. 1177 and 1178, comprises too small a number to permit drawing any conclusions, but apparently offers no particular advantages (see Table 22).

13. GROUP M-COBALT STEELS

The steels in this group (Table 21) are all from the comparison series. All except No. 15, which contained molybdenum in addition, showed high tensile strength, with good ductility and toughness. It is probable that cobalt acts similarly to copper in replacing some of the nickel.

TABLE 11.—Group A—Silicon Steels LESS THAN 1 PER CENT SILICON

	I		Ft 1bs./in.*	128			625	460	246	218	109	121
	ness eral	Sclero- scope	30	28	56		18	19	25	31	29	33
	Hardness numeral	Brinell Sclero-scope	195		187		185	207	188	235	418	364
pe	Reduc-		P. ct.	30.6	26.3		51.5	52.0	13.1	53.8	15.9	4.8
Heat treated		in 2 inches	P. ct.	9.5	11.0		12.5	17.0	8.0	15.0	4.5	1.0
H	Ultimate	strength	Lbs./in. ²	111 200	126 000		99 700	105 000	156 500	121 900	259 400	230 600
	Yield	point	Lbs./in. ² Lbs./in. ²		26 000		57 400		75 000	85 400		
	Propor-	limit	Lbs./in.²	42 000	30 000	COIN		40 000	40 000	45 000	137 000	105 000
	ness	Sclero- scope	28	24	27	IT SILI	78	28	26	28	24	19
	Hardness numeral	Brinell Sclero-	160	163	178	ER CEN	178	196	190	209	241	269
	Reduc-	area	P. ct.	50.2	46.0	MORE THAN 1 PER CENT SILICON	59.0	59.0	51.5	48.1	51.5	45.1
Normalized	Elon-		P. ct.	26.3	25.0	RE TH	22. 5	21.5	23.0	19.0	20.0	15.0
Norm	Ultimate	strength	Lbs./in. ³	89 600	95 400	MO	87 300	94 100	102 900	105 300	122 500	133 400
1	Yield	point	Lbs./in. ²		61 500		61 200	68 500	60 400	006 69	85 900	008 06
	Propor-	limit	P. ct. P. ct. Lbs./in. ²	43 000	52 000		28 000		40 000	62 000	79 000	000 59
	Ē	3	P. ct.		.03				0.06			
Hon		₹	P. ct.	. 45	.00		0.01		8. 6	.02	.0	.01
Composition	- 1	K	P. ct.	. 52	. 58		0.71	.75	.65		.86	.74
ပိ		75	P. ct.	.46	99•		1.10	1.10	1.15			1.20
		ပ	P. ct. P. ct. P. ct. 1	.38	• 39		0.24	. 27	.36	.41		9.
	No.		1269		1104		1198	1199	1102	1201	1164	1200

TABLE 12.—Group B-Nickel-Silicon Steels

-	4
G	1
MILL	3
E	1
۴	4
F	4
THE	7
þ	1
ζ)
ρ	4
COC	1
P	4

		Impact	Ft	340	133	79	110	19	46	92	
	Hardness numeral	Brinell Sclero-		20	34	35	33	30	49	74	
	Har	Brinell		255	286	321	418	340	009	652	
eated	Reduc-	area	D.	39.6	21.1	9.2	7.3	16.0	9.	2.0	
Heat treated	Ultimate Elonga Reduc-	inches	D.	7.0	7.5	3.5	3.0	4.0	1.0	2.0	
	Ultimate	strength	Lbs./in.²	127 000	158 100	294 200	261 600	316 700	192 800	322 700	
	Yield	point	Lbs./in.2 Lbs./in.2 Lbs./in.2	84 900	108 400					262 600	
	Propor-	rional limit	L.bs./in.²	52 000	26 000	155 000	132 000	170 000		154 000	
	Hardness numeral	Sclero- scope		19	31	36	32	28	38	50	
	Hardnes	Brinell		191	217	228	240	418	286	322	
	Reduc-	area	P. P.	54.7	54.4	50.4	54.4	45.2	7.9	20.2	
Normalized	Elonga-	non in z inches	D.		21.5	19.5	21.0	17.5	4.0	10.5	
No	Ultimate Elonga Reduc-	strength	P. ct. P. ct. Ths./in.2 Ths./in.2	94 300	109 300	130 200	121 400	131 400	115 100	172 500	
	Yield		Lbs./in.2	70 700	009 44	91 900	88 600	84 800		125 000	
	Propor-	timit	T.bs./in.2	56 000		71 000	81 000	49 000		123 000	
	i	4	p.		0.27				. 44		
		₹	p.	0.02	0.02	10.	5 Tr.	5 .01	N.D.	Tr.	_
Composition		Z 	p.	0.95 0.61 2.00	0 2.10	6 2.05	0 2.05	5 2.15	0 2.15	4 2.20	_
Com		Mn	p	0.0	. 90	97.	03.	5 . 75	06.	. 94	_
		<u> </u>	l a	5 0.95	0 1.23	1 1.50	2 1.35	4 1.25	6 1.00	7 1.85	_
		ပ	6	0.2	. 40	. 41	.42	. 54	.56	. 57	_
	No.			1202 0. 25 0. 95	1174	1165	1166	1167	1246	1245	

	88	33			
52	55		36	78	
537	512		444	375	
2.7	14.1		1.1		
2.5	5.5		9.	1.0	
286 000	275 900	(a)	286 000	194 500	
	155 000 185 750	(a)			
126 000	155 000		170 000	63 000	
30	32		35		
302	255		241		
4.0	30.0		18.9		
4.5	15.0		11.3		
178 500	136 200		. 117 200		
	80 900				_
	26 000		.03 67 100		
	09.0	:			
0.01	90.	:	.02	Tr.	
3.15	3.00	3, 25	3, 25	3, 20	
0.97	.81 3.00	.90 .80 3.	.65	1.10	
1.00	.65	06.	.52	.51 1.00 1.10 3.2	1
0.44	. 45	. 48	. 49	.51	1
1120 0.44 1.00 0.97	1242 45 .65	2848	2114 49 .52 .65 3.2	1118	

50 50

3 TO 3.25 PER CENT NICKEL, MORE THAN 1 PER CENT SILICON

134	66		88			37	84	100	74	911	83	75	:	112	129	95	20	61	88	61	38	02
	_			-			_	_	_	_											_	
49	29	37	47	51	53	89	46	36	52	45	69	59	32	33	44	46	56	46	37		53	81
382	364	402	564	488	512	287	578	205	555	564	555	555	387	444	546	564	591	532	564		009	627
37.2	15.4	22.9	36.6	25.5	26. 1	2.0	31.3	38.5	38.9	35.6	32.9	1.6	22.8	14.9	34.8	11.1	12.9	24.9	20.3		2.0	
0.9	6.5	8.0	10.0	7.5	15.0	1.0	9.5	10.5	8.5	8.5	9.5	1.0	6.5	0.9	11.0	5.2	7.5	8.0	7.0		.5	
						286 700												312 600			206 500	
	-	-	231 000		000		300	100	200	000	400	:	:	006	000	009	200	100	100	\div		
116 000	134 000	000	000	140 000	000	102 000	000	000	000	000		<u> </u>	000		-		000	:	160 000		130 000	
						57 1								•	•			•		37	27	46
228	444	241	268	249	241	450		569	566	368	588	255	300	255	277	271	592	317	586	586	302	302
54.1	48.3	27.1	45.1	20.0	8.6	3.4		30.3	52.0	9.8	42.3	47.3	11.5	34, 4	48.4	47.3	53, 3	40.9	42.9	41.3	30.6	30.2
23.0	19.0	13.1	19.0	4.5	8.0	1.5		11.0	18.5	8 0	14.0	19.5	4.0	17.5	20.0	17.5	21.0	14.5	15.5	16.5	12.5	5.5
						191 100															148 200	
						83 000		107 500	000 46	145 200	006 96	96 150	:	122 200	000 06	93 600	92 300	108 000			102 700	
49 000	44 000	40 000	39 000	20 000	72 500	41 000	-		52 000		45 000	000 94	000 09		45 000	65 000	23 000	72 000	20 000	71 000	22 000	49 000
	:	-	•		0.33			Tr.				:	:	. 45	:	10.		:	. 45			
0.01	10.	10.	Tr.	.01	-02	.01	Tr.	.02	10.	Tr.	Tr.	Tr.	.33	.12	. 23	. 02	Tr.	Tr.	.17	Tr.	.01	Tr.
						3.15																
0.78	.77	. 85	.77	98.	.90	1.46	1.99	. 86	. 84	1.10	06.	. 82	. 81	. 82	.81	.71	. 78	. 94	- 80	. 79	.97	1.13
1.35	1.35	1.15	1.40	1.10	1.20	1.25	1.25	1.45	1.45	1.45	1.60	1.35	1.05	1.25	1.25	1.35	1.30	2. 20	1.10	1.25	1.25	1. 20
0.26	.35	.36	.39	.40	.40	.40	.40		.40	9.	.40	. 42	. 43	. 43	. 45	. 48	. 49	. 49	. 52	. 52	. 53	
1711	1168	1129	1214	1128	1130	1251	1238	1204	1227	1236	1226	1169	1147	1216	1206	1205	1215	1237	1217	1170	1208	1209

3.25 TO 3.50 PER CENT NICKEL

	Ξ	V)			
	42		42		
	387	-	209	-	
_	32.6		.7		
-	5.5	(a)	.25		
	224 100	(g)	235 000		
	121 000				
-	33		43		
	228		284		
	54.2		6.5 11.6		
	23.5		6.5		
	112 600		132 400		
	Tr 44 000 77 100 112 600		74 000 90 000 132 400		
	44 000		74 000		
			0.03 0.03		
	Tr.		0,03	Tr.	
	3.30	3.49	3,55	3, 50	
	_	1.01	.76	96	
	0.24 1.30 0.76	1.65	.55 1.15	1.50	
	0.24	.45 1.65		.58 1.50	
	172	3	.113	239.	

a Broke in shoulder.

TABLE 13.-Group C-Silicon Steels with Zirconium

		Impact	Ft	10s./in.² 237	352	116	124	82		126	:				
	ness	Sclero- scope		22	82	27	31	35	25	31	54	27	28	44	43
	Hardness	Brinell		187	387	228	187	586	196	569	209	241	262	520	454
ated	Reduc-	area		F. ct. 55.0	54.7	30.1	26.8	19.8	21.3	32.0	4.5	8.5	2.2	2.0	0.
Heat treated	Elonga-	inches		P. ct.	20.5	0.6	12.5	4.0	9.5	1.5	1.5	5.0	.5	1.5	. 25
	Ultimate	strength inches		Lbs./in. ² 99 140	97 100	135 600	140 400	192 600	99 500	148 800	236 000	164 500	147 500	262 000	202 750
		point		Lbs./in. ² Lbs./in. ² Lbs./in. ³ 23 000 62 400 99 140	54 800		91 900		:	128 300	206 000				
	Propor-	limit		Lbs./in. ²	36 000	35 000 -		000 06	24 000	75 000	126 000	54 000	50 000	80 000	
		Sclero- scope		24	11	27	30	17	27	53	52	27	31	62	27
	Hardness	Brinell		179	179	189	202	506	179	212	186	185	202	877	197
	Reduc-	area		P. ct. 60.8	62.9	51.5	53.7	53.0	22.8	52.3	20.3	39.8	22.7	38.6	18.7
Normalized	Slonga-	inches		P. ct.	28.0	22.5	24.5	24.5	10.0	19.0	8.0	19.5	16.0	16.0	11.0
No	Ultimate	strength		Lbs./in. ²	92 200	92 500	100 700	106 800	95 000	115 400	101 000	000 66	110 800	115 400	100 700
		point		Lbs./in.2 Lbs./in.2 Lbs./in.3 28 000 45 000 79 300	63 500	48 000	62 100		50 500	76 500	53 250				26 000
	Propor-	limit		Cbs./in.2 28 000	20 000	32 000		43 000 -	30 000	47 000	39 000	42 000 _	31 000 [.	53 000 .	20 000
	1	17		P. ct. 1	.50	. 50	. 60	. 22	. 20	.03	.15	.03	.11	. 10	60.
13	į	=		P. ct.	T .08 B .05	.10	.07	90.	.03	.03	.01	.02	.02	.04	.02
Composition	:	₹		P. ct.	Tr	Tr.		Tr.	0.05	ij.	.13	.02	.15	. 09	.07
Comp	1	u Wi		P. ct.	.71	.63	.69	.77	. 50	.76	. 55	.67	.78	. 80	.75
		ภี		1.30	1.35	1.55	1.70	1.70	.73	1.55	. 44	.50	. 85	1.15	.54
		ပ		P. ct	. 26	. 33	. 34	.36	.37	. 42	. 42	. 45	. 47	. 51	. 56
	No.			1184 0.23 1.30 0.61	1185	1180	1181	1183	1107	1182	1106	1103	1109	1101	1105

TABLE 14.—Group D.—Nickel-Silicon Steels with Zirconium 2 PER CENT NICKEL

		1	TIMbacı	Ġ	lbs./in. ² 163	202	122	236	101	120	136	81		108	91	88	51	-
			Sclero- scope		-	24	47	32	34	33	32	43	44	37	43	35	37	-
		Hardness numeral	Brinell		418	255	382	241	306	364	289	340	390	47.7	385	444	351	
fod	nen	Reduc-	area		P. ct.	40.3	39, 3	41.3	11.7	24.0	38. 2	23.4	32.2	24.1	31.9	11.7	10.7	
Heat treated	ical ilea		in 2 inches		P. ct.	10.0	4.0	0.6	2.5	2.0	16.5	7.0	4.0	7.0	7.9	5.5	33. 5	-
	•	Ulti-	н		Lbs./in. ² 206 800	125 300	180 500	133 700	201 200	180 200	143 400	175 300	247 500	293 700	283 800	282 100	245 600	
		Yield	point		Lbs./in. ² Lbs./in. ² Lbs./in. ³ 90 000 206 800	000 68	148 900		172 800	148 500	127 300	009 86			254 200		184 700	-
		Propor-	limit		.bs./in.2 L	35 000	67 000 1	46 000	-	-	39 000 1	:	123 000	166 000	146 200 2	126 000	73 000 1	-
			Sclero- scope		25 L	92	31	28	56	25	30	36	21 1	31 10	1	32 1	21	
		Hardness	Brinell Sc		202	212	194	219	526	217	216	286	228	255	:	592	288	-
		Reduc-			P. ct.	57.3	56.1	54.3	54.7	49.3	54.7	50.1	46.0	44.9		42.3	32.7	-
Normalized	danteed		inches		P. ct.	19.5	28.5	23.0	21.5	22. 5	22.0	21.0	21.0	19.5		24.5	18. 5	-
Non	TAOLE	Ulti-				95 600	96 200	006 46	106 800	106 100	103 000	116 600	115 900	139 600		127 600	113 200	-
		Yield	point		Lbs./in. ² Lbs./in. ² Lbs./in. ² 35 000 110 000	000 29	68 400	000 44	77 100 1	77 400 1	75 500 1	87 300 1	78 600 1	86 900 1		-	70 000 1	-
		Propor-			Cbs./in. ² Lt 35 000	30 000	42 000	49 000	34 000	:	000 19	21 000	73 000	000 19	:	49 000 6+	34 000	-
		<u> </u>			o. ct.	T. 80 B. 35	T.38 4	.30	.40	. 45	09.	. 55 5	. 08 7	.13 6	-	. 10 4	. 10	-
		. Ē	=		ct. P.ct. P.ct. F 00 Tr 0.06	.15	0.	. 20	60.	.07	.04	. 04	.01	11.	:	.15	.01	-
200	HOIT	7	₹	=	P. ct.	Tr	Tr	Tr	Tr		0.07	.14	90.	.17		. 12	90.	
Composition	m posi	ž	Ę		P. ct.	2, 05	2.00	2.05	2.10	2, 15	1.75	1.60	2.15	2.00	2, 16	2.10	2. 10	
2	3	3			P. ct.	.76	.67	.67	.77	.70	.62	.76	.75	.87	.71	.97	99.	
		5	กี		P. ct. P. ct. P. ct. 1.35 0.74 2.00	1.50	1.40	1.45	1.40	1.45	. 46	2,00	1.20	1.10	1.26	1.45	1.25	
		()		P. ct.	92.	. 29	. 29	.32	.33	.35	.35	. 38	. 43	. 43	.45	.51	
		No.			1186.	1192	1187	1193	1191	1190	1249	1248	1161	1176	27	1175	1162	-

TABLE 14.—Group D—Nickel-Silicon Steels with Zirconium—Continued 3 PER CENT NICKEL, LESS THAN 1 PER CENT SILICON

									The same (services a response a r													1
			ŭ	Compositio	tion					No	Normalized							Heat treated	ated			
No.								Propor-	Vield	-jiji	Elon-	Reduc-	Hardness	less	Propor-		-HIL		Reduc-	Hardness numeral		Irranart
	O	ii.	Min	Ž	E .	F	72	tional limit		mate strength	in 2 inches	tion of area	Brinell	Sclero- scope	limit	point	strength	in 2 inches	area	Brinell	Sclero- scope	mpact
				1		1	1		0 1		1	1 4			The ling	The fin 2	T be /in 2	į p	Į į			Ft
1222	P. C.	P. Ct.	P. Ct.	3.00	0.04	04 0.01	0.04	ros./m.	LDS./III.2 LDS./III.2	95 700	22. 5	47.3	183	25	137 000	223 800	239 700	8.5	32.4	387	55	91
1212	34	. 80	. 65	3,05	Ä	9	4	42 000	65 500	106 000	24.0	41.7	207	30	100 000	212 500	221 500	10.5	35.6	450	26	100
1223	. 36	.30	. 57	3.00	.03		.07		73 600	106 200	20.0	52.2	196	56	136 000	243 600	269 700	8.5	29. 5	324	36	79
1233	.36	. 27	. 57	3.05	. 02		.16		71 400	103 000	21.5	53.4	265	28		207 600	245 300	8.5	31.9	200	57	74
1240	.37	.65	.70	3.00	. 02	.01	. 12		92 400	156 900	6.5	23.8	566		137 000	203 300	273 700	8.5	28.9	512	29	120
1224	.38	.95	. 45	3.05	Ħ	. 10	. 20	000 09	83 700	106 800	24.0	50.5	418	20	63 000	146 800	175 000	8.5	14.8	340	92	101
1243		.95	. 85	3.00	.01			38 000	173 000	200 600	6.5	16.0	324	30	146 000	249 100	281 800	10.0	38.3	246	22	138
1213		.80	. 65	3.15	T	9.	. 33	42 000	73 000	112 300	20.5	40.9	241	22	150 000	008 602	276 500	0.9	20.8	246	41	38
1232		. 20	. 48	3.00	.01			38 000	007 79	100 300	22.0	48.3	199	27	141 000	228 200	256 100	8.5	32. 2	220	63	74
1115	. 43	. 20	.73	3.15	Tr		. 05	72 000		121 000	16.5	40.2	210	92	62 000		170 000	2.0	11.6	354	32	
1231	. 43	.76	. 81	3.00	.05		.15	59 000	148 900	188 900	3.0	18.4	569	34		211 600	290 100	9.5	36.7	246	48	109
1119	4		.92	3.15	.01	.01	.05	53 000	100 000	198 000	2.5	33, 2	321	35	120 000		297 000	7.0	18.5	246	56	
1158	44.	.75	. 73	3. 10	.16	.03	. 21	47 000	138 200	140 200	8.0	23. 2	447	25	172 000		287 000	0.9	17.8	387	42	46
1111	. 46	. 27	. 59	3.15	.03	.01	.03	77 000	82 200	110 500	17.5	37.3	216	30	95 700		271 000	.7	1.2	438	37	:
26	. 48	.93	96.	3.01	:										(g)	(a)				<u>:</u>	:	27
24	. 48	.57	. 81	3.18											150 000	226 800	281 500	9.0	33. 4	278	99	96
20	. 54	.95	. 72	3, 19											121 700	236 900	301 800	5.7	14.9	545	49	75
1241	.61	.67	.75	3.05	.01	.02	. 10	46 000	112 100	145 800	10.0	37.7	569	33		260 300	290 300	3.0	2.7	591	79	58
1230	1.53	. 22	. 70	3.00	.04	.04	.08	81 000	100 300	151 400	5.0	11.0	298	42						009	73	9 !
								3.1	PER CENT	IT NICK	EL, MC	ORE TH	CAN 1 P	ER CE	NICKEL, MORE THAN 1 PER CENT SILICON	ICON						
1189 0.27	0.27	1.50	0.73	3.15	-	0.04	0.20	44 000		116 100	20.0	44. 4	241	33	110 000	216 000	221 600	18.0	26.8	321	24	109
1188.	118827		. 74			Tr06	. 32			74 300 100 400	24.5	55.3	197	24	80 000	80 000 189 600	199 400	10.0	40.9	344	56	136
		-				-				_												

Burgess Woodward]		Zirconia	um Steels		
183 116 102 130	141 146 117 82 121	88 122 116	112	75 181 95 88	112
39 44 39	22 27 27 48 47 62	34 46	47 50 39 43	55 45 55 69	55 68
382 382 382 488 555	436 248 351 420 319 424	375 53 2 532	454 436 502 512 555	523 387 534 396 555	532 600 555
42.1 27.5 3.9 36.9 34.3 13.6	35. 3 33. 6 36. 5 12. 9 28. 2 28. 2	34.0	26.5 21.7 36.9 4.5 8.4	37.3 37.3 19.8 14.1 12.2	9.9
7.0 3.0 2.0 7.5 8.0	9.6 12.0 6.5 5.5 10.0	6.0	8.5 7.5 10.0 1.0 4.0	7.5 11.3 2.5 8.5 6.0	3.5
	228 800 152 800 232 200 232 000 173 600 252 500		284 800 307 600 242 000 267 500 290 600	257 100 257 100 219 800 325 400 261 600 259 000	300 800 296 800 303 600 280 200
0000	9000 9000 3000 2000 2000 8000 2000		000	300 2 300 2 800 2 800 2 800 2 800 800 800 800 8	
142 189 217 221	202 129 218 216 130 245	153	239	188 199 268 217 217 237	245
	111 000 106 000 90 000 62 000 152 500	130 000	125.000 130 000 136 000 135 000	70 000 1175 000 1110 000	144 100 150 000
33 33 33 33 33	32 38 32 32	33 28	30 32 44 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	38 88	34 33
216 228 248 223 279 279	217 234 307 228	302	268 386 255 375 364	255 275 277	241
53. 0 52. 8 15. 6 52. 3 10. 5	55. 6 44. 7 20. 8 50. 6	52. 0 50. 6 53. 3	22.6 5.2 32.7 5.2 4.2	43.8	13.4 32.2 27.5 1.4
27.5 22.5 12.0 23.5 9.0	24. 0 22. 0 16. 5 24. 0	23.0	20.0	8.0 19.5 5.0 17.0	5.5 14.0 11.0
110 300 109 800 111 100 109 200 140 400 201 500	100 400 107 900 172 100 110 800		220 200 220 200 132 300 259 000 221 000		207 100 143 300 100 600 230 700
67 700 79 000 84 900 82 000 119 000	64 900 78 900 150 300 74 000		142 800	79 400	93 400
000 000 000 000	: 000 000 :	: 00 :	000000000000000000000000000000000000000	000 000	0 0 00
.60 38 .33 52 .50 46 .55 68 .80 } .35 }	70 48 55 60 40 60 16 44,	·	31 59 10 26 12 54 20 50		05 35 05 05 08 50 07 76
FB		T B			
.00 .07 .07 .00 .00 .00 .00 .00 .00 .00			2 2 2 3 3 3		1. 10
10.02 17.17.17.	.02 Tr. .14	. 07			10. Tr. 090
જું છે છે છે છે	2.60 3.10 3.05 3.00 3.10	3.05	3.20 3.10 3.10 3.20		3.20 3.10 2.95 2.74 2.74 3.20
	. 82 . 38 . 57 . 86 . 61	. 38			.84 .66 .93 .93
1. 45 1. 50 1. 50 1. 50 1. 10 1. 30	1.35 1.35 1.10 2.00 1.00 1.01	1. 10		1. 20 1. 65 1. 05 1. 56 1. 55 1. 85	1.05 1.10 1.05 1.00
	8. 8. 8. 8. 8. 8. 8. 8. 8. 8. 8. 8. 8. 8	.36	. 38 . 39 . 88 . 88 . 89 . 89		.45 .46 .46
1235 1197 1196 1211	2 1219 1210 1290 21	1194 1218 1225	1293 1144 1157 1131	1145 1220 14 1289	1146 1138 1112 25

a Broke in shoulder.

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TABLE 14.—Group D—Nickel-Silicon Steels with Zirconium—Continued 3 PER CENT NICKEL, MORE THAN 1 PER CENT SILCON—Continued

	Impact		Ft Ibs./in.² 75	75		54		102	168	145	:		41	124	
	ness	Sclero- scope	25	50	36			4	20	62	19	55	62	4	
	Hardness	Brinell Sclero-	491	402	440			320	324	440	584	512	, 564	555	
reated	Reduc-	area	P. ct.	1.3	4.0			46.3	28.6	29.0	18.3	14.8	4.7	32.0	
Heat treated	Elon-		P. ct.	1.5	3.0			8.	7.8	8. 6 (a)	8.5	7.0	1.0	11.5	
	- diti-	strength	Lbs./in. ² 234 300	286 100 216 800	291 200		CICON	156 600	208 700	245 000	282 500	292 000	286 800	253 900	
	Yield		Lbs./in. ² Lbs./in. ² Lbs./in. ² 166 000 234 300				MORE THAN 3.25 PER CENT NICKEL AND MORE THAN 1 PER CENT SILICON	142 000	187 800	202 800	254 700		226 200	259 400	
	Propor-	limit	.bs./in. ²	150 000	118 000		1 PER C		110 000	125 800 (a)	8	136 000	130 000	112 500	
			32	35	20	35	THAN	41				33			
	Hardness	Brinell Sclero-	266	364	255	321	MORE '	286				292		-	
	Reduc-	area	P. ct.	24.3	11.6	21.9	AND	49.3				31.9			
Normalized		in 2 inches	P. ct.	18.5		8.0	NICKEI	15.5				12.0			
Noi	Ulti-	mate strength	Lbs./in. ² 155 700	154 800	195 200	154 300	CENT	133 400				148 500			
	Yield	point	Lbs./in.² Lbs./in.² Lbs./in.²	106 800		106 600	3.25 PER	006 96							
	Propor-	tional	Lbs./in.2	57 000	000 09	000 89	THAN	39 000				46 500			
		17	P. ct.	90.	. 25	T.85 B.71	MORE	T .65				T.70 B.30			
	i	F	.ct. P.ct.	.04		90.		0.03				. 02	:	:	
tion		₹	1 40	10.		10.		0.01				.01	:		
Composition		Z	P. ct.	3.15	3.00	2.90		3, 55	7.86	3.64	3.33	3.30	3, 35	3.40	5.10
ပိ		Mn	P.ct.	. 93	.72	. 85		0.95	.36	1.14	.77	.75	. 78	1.01	3
		ii.	P. ct.	1.78 .93	1.20	2.45		1.55	. 47	2.16	1.82	1.65	1.59	1.56	1.10
1		ပ	P.ct.	.48	.51	. 54			. 19	.37		.39	.39	. 41	. 10
	No.		P.ct.	129248	1159	1247		1250 0.15	:::	8	7	1133	16	18	

a Not tested in heat-treated condition.

TABLE 15.—Group E—Cerium Steels WITHOUT NICKEL

		Impace	Ft Ibs./in. ³ 72 128		72	174	71	252	89	47	33
		Sclero- scope	32		52	58	57	51			52
	Hardness	Brinell	418		555	555	228	009			530
ated	Reduc-	of area	P. ct. 0.7 10.0		37.2	7.2	10.4	11.0	2.1		14.1
Heat treated	Elonga-	non in 2 inches	P. ct.		8.5	7.5	8.6	5.5	2.5		1.0
	Ultimate Elonga-Reduc-	strength 2 inches of area	Lbs./in.² 128 200 188 900		311 100	298 000	202 500	324 800	177 500	(a)	251 700
	Yield	point	Lbs./in. ² Lbs./in. ² Lbs./in. ² 85 000 128 200 120 000 188 900		206 500	244 900	149 000	288 900	154 500	(a)	
	Propor-	limit	Lbs./in.² 85 000 120 000			175 000	65 000	83 000	91 000		140 000
	ness eral	Sclero- scope	17 25	-	24	39	37	36	34	48	40
	Hardness	Brinell	163	CKEL	269	285	217	302	187	359	396
	Reduc-	of area	P. ct. 5.4 31.1	WITH NICKEL	40.8	37.6	17.8	17.3	16.0	36.1	
Normalized	Elonga-	uon in 2 inches	P. ct. 6.3 8.5	A	15.6	7.5	21.0	5,5	7.6	10.5	1.0
No	Ultimate	strength 2 inches of area	Lbs./in.² 93 600 84 700		143 900	143 400	127 300	158 600	109 100	171 000	154 300
	Yield	point	Lbs./in.² 56 600 71 200		134 300	102 200	80 000	142 600	74 000	115 300	
	Propor-	limit	Lbs./in.² Lbs./in.² Lbs./in.² 37 000 56 600 93 600 27 000 71 200 84 700			60 000	53 000		20 000	100 000	70 000
	d	ي	P. ct. 0. 20 . 35		0.01	90.	T.31	T.55 B.35	T1.35 B.66	T.22 B.07	Cu. 62
tion		Ę	P. ct.		2.95	2.80	2.95	3.00	2.65	2, 25	2.90
Composition	le p	ши	P. ct. 0.68 .69		0.71	.73	1.15	.91	.90	.82	1.04
0	5	70	P. ct. P. ct. 0.39 0.75	-	1.30	1.70		1.30	. 25	1.25	1.35
	(ט	P. ct. 0.39		0.45	.41		. 44	.39	. 74	.51
	No.		12681272.		1260	1256	1259	1252	1258	1253	1281

a Broke in shoulder.

TABLE 16.—Group F-Copper Steels

			0	Composi	sition						No	Normalized							Heat treated	ated			
° Z						į	t		Propor-		Ultimate	Elonga-			Hardness numeral	Propor-	-	Ultimate	Elonga		Наго	Hardness	
	ပ	22	MM m	į į	TA T	=	17	5	tional limit	point	strength	tion in 2 inches	area area	Bri- nell	Sclero- scope	limit	point	strength	tion in 2 inches	fron or area	Brt- nell	Sclero- scope	Impact
	P. ct. P. ct. P. ct.	45	, c	P. ct	P. ct.	P. ct.	P. C.	ا ن ن	Cbs./in.	ct. P. ct. P. ct. Lbs./in. ² Lbs./in. ² Lbs./in. ²	Lbs./in.2					Lbs./in.2	Lbs./in.	Lbs./in. ² Lbs./in. ² Lbs./in. ² P. ct.	P. C.	P. ct.			Ft
:	1285 0.35 1.40	.40 0	0.76	2.55 0	0.05	Tr.	0.70	0.02 Tr. 0.70 0.62	43 000	133 100	152 700		23.8	286	53		233 300	248 000	7.5		402	55	83
128245	. 45	1.10	.84	1.90	Tr.		1.35	1.35	80 000	128 200	150 800	11.0	27.2	320	37		273 400	313 300	7.5	36.6	546	57	28
123646	.46	1.30	. 82	2.55	Tr.		:	.64	70 000	134 300	143 700	12.0	38.1	292	38	186 000	279 100	327 900	4.5	7.9	555	63	99
128049	. 49 1	. 25 1	1.03	2.45	.01	:	:	.55	55 000	171 800	173 600	0.9	14.8	326	35	150 000		202 2002			550	09	20
1283	.50 1	1.25	. 78	2.60	.01		:	.36	85 000	131 500	147 500	12.0	44.7	285	38		261 600	297 900	1.0	1.9	573	68	25
1281	.51	. 35	1.04	2.90		:		.62	20 000		154 300	1.0		396	40	140 000		251 700	1.0	14.1	230	52	33
1279 58		. 23	96.	2.45	10.	:	.01	-62	71 000	131 500	139 700	11.5	19.7	279	37	140 000		251 700?		.7	642	70	34

TABLE 17.—Group G—Boron Steels WITHOUT NICKEL

1141													
	10000	Impac	Ft lbs./in.ª	37	48	36		161	48	46	109	35	41
	Hardness numeral	Sclero- scope		30	43	52		38		52	09	55	89
		Brinell		217	555	387		387	380	387	470	425	009
ated	Reduc-	area	P. ct.	19.1	9.8	.7		2.7	2.0	5.9	7.9	7.9	3,5
Heat treated	Elonga-	inches	P. ct.	3.9	.5	-:		0.8	3.2	4.3	2.3	3.0	3.0
	Ultimate Elonga- Reduc-	strength	Lbs./in.²	96 100	150 000	85 400		170 900	208 300	214 500	226 700	256 300	323 100
		point	Lbs./in. ² Lbs./in. ² Lbs./in. ²		147 400					169 700	205 100	244 900	
	Propor-	limit	Lbs./in.²	51 000	81 000			114 000	112 000	113 000	135 000	100 000	170 000
	62	Sclero- scope		15		83		22	21		24	35	34
		Brinell		156	217	187	CKEL	220	196	175	202	292	285
	Reduc-	area	P. ct.	23.8	25.6	41.7	WITH NICKEL	10.6	5.4	20.8	46.8	31.1	22.6
Normalized	Elonga-	inches	P. ct.	10.7	11.5	17.0	W	10.5	14.0	10.8	12.0	17.0	11.0
ğ	Ultimate Elonga- Reduc-	strength	Lbs./in.²	69 800	84 600	100 500		110 400	65 800	90 700	160 900	120 600	139 900
	1		P. ct. P. ct. Lbs./in.2 Lbs./in.2 Lbs./in.2	46 300	28 000	84 800				65 000			132 900
	Propor-	limit	Cbs./in.²		54 000			40 000	48 000	55 000	53 000	20 000	27 000
	F	a -	P. ct.	0.39	.57	90.		0.09	. 30	.50	. 10	. 10	. 08
		₹	P. ct.	0.00	.02	.03		0.02	. 02	Tr.		.02	.01
Composition	;	Į.	P. ct. P. ct. P. ct.					3.00	3.00	3.05	3.55	2.80	2, 90
Comp		u W	P. ct.	0.68	.80	69.		0.84	.77	. 67	. 58	. 67	. 50
	į	ñ	P. ct.	0.24	1.50	. 33		1.20	1.30	. 41	.36	1.25	.36
		ပ	P. ct.	0.16	. 21	.45		0.16	.18	. 19	. 26	. 47	69.
	No.			1267 0.16	1261	1274		1278	1263	1264	1277	1276	1275

38 34 34

53

556 556 540

1.0

5.1

TABLE 18.—Group I-Molybdenum Steels

NICKEL-MOLYBDENUM

		ımbacı	Ft Ibs./in.2	168			72			-=	
	Hardness numeral	Sclero- scope		20	33	36	70	69	_		
		Brinell Sclero-		324	444	393	509	589			
eated	Reduc-	area	P. ct.		8.4	12.2	(a)	26.2			
Heat treated	Elonga	inches			5.5	6.5	(g)	7.8			
	Ultimate Elonga - Reduc-	strength	Lbs./in.	208 700	300 100	257 700	226 700?	288 600			
	Yield	point	Lbs./in.2 Lbs./in.2 Lbs./in.2	187 800				229 500			
	Propor-	limit	Lbs./in.²	110 000	83 000	100 000	140 000			IMI	
		Sclero- scope			35	37				ANADIT	
	Hardness numeral	Brinell Sclero-scope			286	302				CHROME-MOLYBDENUM-VANADIUM	
774	Reduc-	area	P. ct.		25.9	20.2				LYBDE	
Normalized	Elonga-	inches	P. ct.		23.0	21.0				ME-MO	
Ä	Ultimate Elonga- Reduc-	strength	P. ct. P. ct. Lbs./in.2 Lbs./in.2 Lbs./in.2 P. ct.		146 000	168 700				CHROI	
	Yield	point	Lbs./in.2								
	Propor-	limit	Lbs./in.2		000 99	37 500					
	Č		P. Ct.				70 0.38	• 59			>
	,		P. ct.	0.29 0.54			.70	.37 .27	_		ర
sition	7,5								_	3	Mo
Composition	2		t. P. ct	5 7.86	3 3.25	1 3.20	3.45	3.06	_		Mn
3			P. C	7 0.36	.83	. 84	8 1.09	9	_		Si
		ة د	P. ct. P. ct. P. ct. P. ct.	0.19 0.47	42 1.4	44 1.8	43 1.2	44 1.4	-		v
	No.		<u> </u>	110	1135 42 1.45	113644 1.80	15 43 1.28	344 1.46			No.

>	0.28
ರ	1.30
Mo	0.84
Mn	0.87
Si	0.22
O	0.34
No.	96

a Broke in shoulder,

TABLE 19.-Group J-Nickel-Chromium Steels

	Import	The second second	Ft Ibs./in.²	113		42	95
	ness	Sclero- scope		57	22	51	72
	Hardness numeral	Brinell Sclero-		532	615	578	567
ated	Reduc-	area	P. ct.	29.5	20.1	11.5	3.3
Heat treated	Elonga-	inches	P. ct.	7.5	. 8	0.9	1.5
1	Ultimate Elonga-Reduc-	strength	Lbs./in.²	258 600		290 400	262 600
		point	L bs./in.2	257 700	247 900		227 800
	Propor-	limit	Lbs./in. ² Lbs./in. ² Lbs./in. ² P. ct.	134 000 257 700 258 600	247 900	153 000	140 000
		Sclero- scope		41		19	
	Hardness	Brinell Sclero-		444		512	
	Reduc-	area	P. ct.	28.2		1.4	
Normalized	Elonga-	fion in 2 inches		7.0		1.5	
No	Ulfimate	strength inches area	Lbs./in.2	201 700		198 900	
	Yield	point	Lbs./in.2	198 400			
	Propor-	tional limit	P. ct. P. ct. P. ct. P. ct. P. ct. Lbs./in. Lbs./in. Lbs./in. P. ct.	50 000 198 400 201 700		113 000	.10 .25 .60
			, c	. 14	. 54		.09
		Zr	P. ct.	07 0.10 1.14	52.	02 1.13	. 25
ion	-	¥	P. ct.	0.02		.02	.10
Composition		Z	P. ct.	3.60	3.54	3.55	3.31
Cor		Mn	P. Ct	0.48 3.60	. 99	2	.95
		<u> </u>	P.	0.16	.39 1.66	. 85	. 43 1.50
		O	D. C.	0.38 0.16		. 43	
	No.			1155	12	115643	19

TABLE 20.—Group K-Vanadium Steels

		rimbaci	Ft	380	121	123	75	91
	Hardness numeral	Sclero- scope		27	43	69	09	40
		Brinell		196		512	627	340
ited	Reduc-	area	P. ct.	45.3	37.6	3.4	7.8	31.0
Heat treated		in 2 inches	P. ct.	11.7	9.0	.1	5.0	2.3
1	Ultimate	strength	Lbs./in.2 Lbs./in.2 Lbs./in.2	115 300	288 400	292 300	343 600	150 900
	Yield	point	Lbs./in.²	95 400		224 600	276 500	
	Propor-	limit	Lbs./in.2	85 000	130 000	170 000		110 000
	ness	Sclero- scope		24	38		56	22
	Hardness numeral	Brinell		208	255		302	217
	Reduc-	area	P. ct.	55.2	46.9		48.4	20.8
Normalized		in 2 inches	P. ct.	22.0	16.0		19.5	7.5
Non	Ultimate	strength	ct. P. ct. Lbs./in. ² Lbs./in. ² Lbs./in. ² P. ct. P. ct.	107 100	145 300		139 000	116 600
	Yield	point	Lbs./in.2	79 300	87 300		103 200	77 500
	Propor-	limit	Lbs./in.²	71 000	51 000		83 000	000 09
	;	>	P. ct.	0,33	.30	. 21	. 32	. 34
ion	ž	Ę	P. ct.		3.00	3.06	3.15	
Composition		11 147	P. ct.	0.46	. 79	1.14	.79	
ပိ	č	<u> </u>	P. ct. P. ct. P. ct. P.	0.45 0.38	1.35	.56 1.09	1.30	
	7	٠	P. ct.	0.45	. 38		9.	. 74
	No.			1273	1173	23	1207	1271

TABLE 21.—Group M—Cobalt Steels

		Timbacc	Ė	lbs./in.2	66	181	72	
	Hardness	Sclero- scope			09	64	70	69
	P-4	Brinell Sclero-			512	534	209	589
eated	Reduc-	area		P. ct.	38.4	19,8	(a)	26.2
Heat treated	Elon-	in 2 inches		P. ct.	8.7	8,5	(a)	7.8
	Ultimate	strength in		Lbs./in.2 Lbs./in.2 Lbs./in.3 P. ct.	277 900	325 400	226 700?	288 600
	Yield	point		Lbs./in.2	143 600 238 100 277 900	175 000 268 300		229 500
		limit		Lbs./in.2	143 600	175 000	140 000	
	Brinell hard-	ness nu- meral						
	Reduc-	area		P. ct.		-		
ized	Elon-	in 2 inches		P. ct.				
Normalized	Ultimate	strength		P. ct. P. ct. P. ct. Lbs./in.2 Lbs./in.2 Lbs./in.2 P. ct.				
	Yield	point		Lbs./in.2				
	Propor-	limit		Lbs./in.2				
		j		P. ct.	1.10	.37	. 38	. 59
	>	OTAT		P.ct.			0.70	.27
ion		13		P. ct.	0.25	.34		.37
Composition	į	E		P.ct.	2. 21	3.01	3.45	3.06
ပိ	2	T T T		P.ct.	0.69	1.08	1.09	06.
	ö	5		P ct.	0,92	1.56	1.28	1.46
	7	ر 		P. ct. P	0.37	• 43	.43	**
	No.				22	14	15	en

a Broke in shoulder.

TABLE 22.—Miscellaneous Steels
CHROMIUM-TUNGSTEN STEELS

	1	_	Ft lbs./in.³	78
		Impact		32
	ness	Sclero- scope		28
111	Hardness	Brinell Sclero-		512
Ped	Re-	tion of area	Lbs./in. ² Lbs./in. ³ Lbs./in. ³ P. ct. P. ct.	11.0
Heat treated	Elon-	Elon- gation in 2 inches		e, S
		Ultimate gation duc- strength in 2 tion of inches area B		58 100
		Yield U		50 150 000 205 000 258 100 8.5 11.0 512
	-			200
	Propo	1d Ultimate gation duc- nt strength in 2 flon of inches area Brinell scope limit		150 00
	iness	Sclero- scope		
	Hard	Brinell		
-	Re-	Re- duc- tion of area		3.0 10.9
Normalized	Elon-	ln 2 inches	P. ct.	3.0
Nor		strength	P. ct. P. ct. 10.10 Cr-1.95, Lbs./in. 2 Lbs./in. 2 Lbs./in. 3 P. ct. P.	W-0.90 Cr-2.00, 66 000 170 900 206 500 W90
	3	point	Lbs./in.²	170 900
	Propor-	Propor- tional limit		000 99
		Ti Other		W-0.90 Cr-2.00, W90
		Ol ele		
g		Zr		1163
Omposition		- F		
Con		Z	P. ct.	3.50
		Mn	P. ct. P. ct. P. ct. P. ct. P. ct. 1777 0.31 0.13 0.41 3.75 0.10	117832 .14 .38 3.50
4		No.		41.
		ပ	P. ct.	. 32
6050	825029 99 A			1178.

		35.2	8.7	3.4	
		10, 5	2.5	1.0	
		309 800	282 800	299 800	
		000 561	191 900		_
		50 000 1	45 000 1	30 000	
	-	35 1	34 1	37 1	_
	-	89	17	90	_
ELS	_	6	33	- A	_
I SIF	_	12.9	80	2.7	
UKANIUM STEELS		0.9	3.0	.5	
UK		183 500	239 700	176 400	
	_	133 200	233 700	169 100	
111		:			
		U-0.34	1229 45 1.05 . 75 3.00 . 01 U— .36 233 700 239 700 3.0 8.5 317 34 145 000 191 900 282 800 2.5 8.7	U52	
,		:	:		
			<u>:</u>		
		0.01	.01	.01	_
	_	0 3.00	5 3.00	4 3.00	
	_	30 0.9	05 .7	20 .8	
	_	43 1.	45 1.	63 1.	1
		12440.	1229	1228	

63593°-22---4

VII. SUMMARY AND CONCLUSION

- 1. About 193 heats of steel, containing in various combinations the principal variable elements of carbon, silicon, nickel, aluminum, titanium, zirconium, cerium, boron, copper, cobalt, uranium, molybdenum, chromium, and tungsten, have been studied.
- 2. None of the steels presented any difficulties in rolling into plate except those containing boron.
- 3. The usual mechanical and impact tests were carried out on all of the steels. It is shown that steel containing 0.40 to 50 per cent carbon, 1 to 1.50 per cent silicon, 3 to 3.25 per cent nickel, and 0.60 to 0.80 manganese and deoxidized with a simple deoxidizer such as alumium can be produced having a tensile strength of approximately 300 000 lbs./in.² with excellent ductility and toughness. This type of steel is recommended for structural material.
- 4. Although the same high properties are obtained in steels of the above composition with the aid of additional elements, it does not appear necessary to resort to such additions of more costly alloying elements.
- 5. Zirconium, like titanium and aluminum, acts primarily as a scavenger, and when it is not removed as part of the slag remains in the steel in the form of square bright-yellow inclusions not directly visible at magnifications lower than 500×. It is not considered that these inclusions can be very beneficial, and if they are segregated into groups and rolled out into thin platelike streaks they may be detrimental.
- 6. Of the other elements that are regarded as special alloying additions, chromium, tungsten, vanadium, and molybdenum go into solution and produce a martensitic pattern in the air-cooled specimens. Cerium and uranium act in a similar manner, but also show characteristic inclusions. Copper goes into solution, but a larger amount is required to produce a martensitic pattern in the air-cooled samples than for the others. Boron forms a complex eutectic, probably that of an iron-carbon-boron compound with iron. This eutectic is fusible at the temperatures ordinarily used in rolling, but at slightly lower temperatures steel containing boron can be rolled successfully. Hot working breaks up the eutectic, and spherical hard particles, similar to iron carbide globules, are formed.

VIII. ACKNOWLEDGMENTS

An investigation covering so many fields of work required the cooperation of many individuals. The authors consider that any merits the investigation may possess are largely due to their collaborators.

The development of the methods of chemical analysis was the contribution of Dr. G. E. F. Lundell, who, assisted by H. B. Knowles and part time by Ensign R. McLane, J. R. Eckman, and Miss E. R. Ward, also made the chemical analyses for the unusual elements. The rolling of the plates was mainly carried out by R. G. Waltenberg, assisted by R. D. France and W. M. Laughton. The last and F. C. Speidel assisted in determining the mechanical properties. Most of the heat treatment was done by H. R. Yerger and the microexaminations entirely by S. Epstein, and H. Scott was responsible for most of the results in thermal analysis.

WASHINGTON, March 30, 1921.

APPENDIX

THE DETERMINATION OF ZIRCONIUM IN STEEL 1

By G. E. F. Lundell and H. B. Knowles

(a) PRELIMINARY STATEMENT

The method developed at the Bureau of Standards permits the determination of silicon, aluminum, titanium, and zirconium in one portion of the steel and provides for the following possible interfering elements: Tungsten, chromium, uranium, cerium, manganese, phosphorus, vanadium, molybdenum, copper, nickel, and cobalt.

(b) METHOD

Dissolve 5.00 g of the steel in 50 cc of hydrochloric acid (sp. gr. 1.2) with gentle warming and the addition of one cc portion of nitric acid from time to time to insure solution of the zirconium and titanium and also oxidation of the iron.

When solution is complete, evaporate to dryness, take up in 10 cc of hydrochloric acid (sp. gr. 1.2), again evaporate to dryness, and finally bake at a gentle heat in order to decompose nitrates. Cool, take up in 50 cc of 1:1 hydrochloric acid, and filter when the iron is completely in solution. Wash the residue with hot 3 per cent hydrochloric acid. Save the filtrate and washings.

Ignite the residue and paper in a platinum crucible, cool, and weigh. Treat with r cc of sulfuric acid (r:r) and sufficient hydrofluoric acid, fume off in the usual manner, ignite and weigh to obtain silica, and calculate silicon. Fuse the slight residue left after the hydrofluoric acid treatment with a small amount of potassium pyrosulfate, dissolve in 10 to 20 cc of 5 per cent sulfuric acid and add the solution to the acid extract from the ether separation obtained as described below.

Evaporate the filtrate and washings from the silica determination to a sirupy consistency, take up in 40 cc of hydrochloric acid (sp. gr. 1.1) and extract with ether in the usual manner. (The ether extract will contain most of the molybdenum, and this element may be qualitatively tested for in it. If molybdenum is present, it is more conveniently determined in a separate portion of steel.) The acid extract will contain some iron and all of the zirconium, titanium, aluminum, nickel, chromium, etc.

Gently boil off the ether in the acid extract, add the matter recovered from the silica, oxidize ferrous iron with a little nitric acid, dilute to 300 cc, cool, and precipitate with 20 per cent sodium hydroxide solution, adding 10 cc in excess. The sodium, hydroxide solution should be as pure as possible and free from carbonate. Filter and save the filtrate. Dissolve the precipitate in warm dilute 1:1 hydrochloric acid, repeat the sodium hydroxide precipitation, filter, and combine the sodium hydroxide filtrates. Dissolve the sodium hydroxide precipitate in warm dilute 1:1 hydrochloric acid and reserve the solution for subsequent analysis.

It is advisable to treat as follows the filter or filters used above: Ignite in platinum, fuse with sodium carbonate, digest the cooled melt with hot water, wash the residue, discard the filtrate and washings, dissolve the residue in hot 1:1 hydrochloric acid, and add to the main acid solution. This precaution makes certain the recovery of any zirconium held back on the filter as zirconium phosphate insoluble in acid.

¹ J. Ind. and Eng. Chem., 12, p. 562; 1920.

Determination of Aluminum

(a) In the absence of chromium and uranium add a few drops of methyl red to the sodium hydroxide filtrate, neutralize with hydrochloric acid, add 4 cc of concentrated hydrochloric acid per 100 cc of solution, boil, make barely alkaline with ammonium hydroxide, continue the boiling for 3 minutes and set the beaker aside for 10 minutes. If no precipitate settles out, the absence of aluminum is assured. If a white precipitate settles out, aluminum is indicated. This precipitate is always contaminated by phosphorus pentoxide and must be purified as follows: Filter without washing, discard the filtrate, and dissolve the precipitate in warm 1:1 hydrochloric acid. Dilute the solution to 50 cc, make alkaline with ammonium hydroxide, neutralize with nitric acid, and add 2 cc in excess. Warm to 50° C, precipitate the phosphoric acid with molybdate reagent in the usual manner, filter, and wash the phosphomolybdate with an ammonium acid sulfate solution. Precipitate the aluminum in the filtrate as directed above, filter without washing, dissolve the precipitate in warm 1:1 hydrochloric acid, reprecipitate, filter, wash slightly with 2 per cent ammonium chloride solution, and ignite in a platinum crucible. The ignited residue is usually contaminated by silica. Therefore a sulfuric acid-hydrofluroic acid treatment, followed by ignition to alumina over the blast lamp, should be performed. (The sodium hydroxide reagent must be tested for substances precipitable by ammonia, and appropriate corrections must be made in the aluminum determination when these are present.)

(b) In steels containing chromium proceed as above until the filtrate from the molybdate precipitation is obtained. Then make the solution ammoniacal, oxidize with a little bromine water, make just acid with 1:2 nitric acid; add ammonium hydroxide in slight excess, heat to boiling, filter, dissolve the precipitate in dilute hydrochloric acid, and reprecipitate the aluminum hydroxide as directed above.

(c) In steels containing uranium the only modification which is required is the substitution of ammonium carbonate for ammonium hydroxide as the final precipitant of the aluminum hydroxide.

(d) In steels containing vanadium, alumina which is obtained by the above procedures from steels containing vanadium is contaminated by this element. When dealing with these steels, proceed as follows: Fuse the weighed residue with pyrosulfate, extract the cooled melt with 5 per cent sulfuric acid, reduce the vanadium in a Jones reductor having ferric alum in the receiver, titrate the reduced solution with standard permanganate, calculate the vanadium as V_2O_5 , and subtract from the original weight.

Determination of Zirconium and Titanium

Dilute the hydrochloric acid solution to 250 cc, neutralize with ammonium hydroxide, so as to leave approximately 5 per cent (by volume) of hydrochloric acid, add 2 g of tartaric acid, and treat with hydrogen sulfide until the iron has been reduced. Filter if the sulfide group is indicated. Make the hydrogen sulfide solution ammoniacal and continue the addition of the gas for five minutes. Filter carefully and wash with dilute ammonium sulfide-ammonium chloride solution. Filter through a new filter if the presence of iron sulfide in the filtrate is indicated. Save the filtrate. (The sulfide precipitate consists of ferrous sulfide in addition to the greater part of any nickel, cobalt, and manganese present in steel. It is preferable to determine these in separate portions of the steel.)

Neutralize the ammonium sulfide filtrate with sulfuric acid, add 30 cc in excess, and dilute with water to 300 cc. Digest on the steam bath until sulfur and sulfides have coagulated, filter, wash with 100 cc of 10 per cent sulfuric acid, and cool the filtrate in ice water. Add slowly and with stirring an excess of a cold 6 per cent water solution of cupferron. (The presence of an excess is shown by the appearance of a white cloud, which disappears, instead of a permanent coagulated precipitate.) After 10 minutes filter on paper, using a cone and very gentle suction, and wash the pre-

cipitate thoroughly with cold 10 per cent hydrochloric acid. Carefully ignite in a tared platinum crucible, completing the ignition over a blast lamp or large Meker burner, cool, and weigh the combined zirconinum and titanium oxides. Fuse with potassium pyrosulfate, dissolve in 50 cc of 10 per cent (by volume) sulfuric acid, and determine titanium colorimetrically or volumetrically. Calculate titanium oxide, subtract the weight found from that of the combined oxides, and calculate zirconium.

(c) NOTES ON THIS METHOD

- r. Phosphorous pentoxide contaminates the precipitate to so slight an extent that it can be disregarded.
- 2. Vanadium interferes no matter what its valency. The interference is not quantitative. If present in the steel, proceed as usual through the weighing of the cupferron precipitate. Then fuse thoroughly with sodium carbonate, cool, extract with water, filter, and determine the vanadium in the filtrate by adding sulfuric acid, reducing through a Jones reductor into a solution of ferric alum-phosphoric acid and then titrating with standard permanganate. Vanadium is thus reduced to V_2O_2 and then oxidized to V_2O_5 . Calculate V_2O_5 and subtract from the combined oxides. Ignite in the original crucible the matter insoluble in water, fuse with potassium pyrosulfate and proceed as directed for titanium.
- 3. Tungsten does not interfere, since it is separated from zirconium and titanium by the sodium hydroxide treatment and from aluminum by the ammonium hydroxide precipitation. If tungsten is present in large amount it may be found desirable to fuse the nonvolatile residue from the silicon determination with sodium carbonate, extract with water, filter, dissolve the residue in hot 1:1 hydrochloric acid, and add to the acid extract from the ether separation.
- 4. Uranium is partially carried down when present in the quadrivalent condition, but not at all in the sexivalent state. If this element is suspected, boil out all hydrogen sulfide before the cupferron precipitation, oxidize with permanganate to a faint pink, cool, and proceed with the cupferron precipitation.
- 5. Thorium and cerium interfere, but they are not thrown down quantitatively. In case these elements are suspected the peroxidized solution used for the titanium determination must be quantitatively preserved and reduced with a little sulfurous acid. The rare earths are then separated by Hillebrand method,² as follows: Precipitate the hydroxides with an excess of potassium hydroxide, decant the liquid, wash with water once or twice by decantation, and then slightly on the filter. Wash the precipitate from the paper into a small platinum dish, treat with hydrofluoric acid, and evaporate nearly to dryness. Take up in 5 cc of 5 per cent (by volume) hydrofluoric acid. If no precipitate is visible, rare earths are absent. If a precipitate is present, collect it on a small filter held by a perforated platinum or rubber cone and wash it with from 5 to 10 cc. of the same acid. Wash the crude rare-earth fluorides into a small platinum dish, burn the paper in platinum, add the ash to the fluorides, and evaporate to dryness with a little sulfuric acid. Dissolve the sulfates in dilute hydrochloric acid, precipitate the rare-earth hydroxides by ammonia, filter, redissolve in hydrochloric acid, evaporate the solution to dryness, and treat the residue with 5 cc of boiling hot 5 per cent oxalic acid. Filter after 15 minutes, collect the oxalates on a small filter, wash with not more than 20 cc of cold 5 per cent oxalic acid, ignite, and weigh as rareearth oxides which are to be deducted from the weight of the cupferron precipitate.

The above procedure does not give an absolutely quantitative recovery of the rare earths. Experiments indicate a recovery of approximately 85 per cent of the rare earths present in residues containing 100 mg of zirconia, 2 mg of thoria, and 2 mg of ceria. Attempts which were made to omit the preliminary separation of the rare earths, as fluorides, were unsuccessful.

² U. S. Geol. Survey, Bul. 700, p. 176.

- 6. Instead of the prescribed treatment for the removal of the bulk of the iron, Johnson's ³ method of fractional precipitation with ammonium hydroxide may be used. When using this method, it is recommended that the 1:1 hydrochloric acid solution of the ammonium hydroxide precipitate should be further treated as given in the Bureau of Standards method beginning with "oxidize * * * and precipitate with a 20 per cent sodium hydroxide solution." In Johnson's procedure silicon must be determined in a separate portion.
- 7. After considering the method and studying the notes the reader might ask the question, "Why not use ammonium hydroxide instead of cupferron as the final precipitant?" The disadvantages of such a procedure are the following: (a) The necessity for destroying the tartaric acid which is in the solution, with attendant danger of contamination by material resulting from the attack on glassware; (b) the coprecipitation of phosphorus and also chromium and uranium when they are present.

The advantages of an ammonia precipitation are: (a) It is a cheaper reagent; (b) the precipitation of cerium would be complete instead of partial.

The following scheme of analysis is now being tested at this Bureau: Zirconium, titanium, aluminum, cerium, chromium, vanadium, etc., are first separated from the bulk of the iron by Johnson's method, and the hydrochloric acid solution of this precipitate is then treated with sodium hydroxide and sodium peroxide as described by Noyes, Bray, and Spear.⁴ It is hoped that this treatment will quantitatively precipitate iron, zirconium, titanium, and cerium, leaving such elements as aluminum uranium, vanadium, chromium, tungsten, molybdenum, and phosphorus in solution. Iron, manganese, and the greater part of the copper, nickel, and cobalt are next separated by precipitation with ammonium sulfide in the presence of tartrate, as recommended by Thornton,⁵ and zirconium, titanium (and cerium) are finally precipitated by ammonia after destroying the tartaric acid. The ignited and weighed precipitate is then treated for titanium and the rare earths as described in the Bureau method.

(d) CONFIRMATORY EXPERIMENTS

Below is given a summary of the data obtained in the analysis of the Bureau of Standards acid-open-hearth steel No. 20a to which definite amounts of standardized solutions were added.

No.	V present G	Cr present G	Cu present G	Ni present G	Al added G	Al found G	T added G	Ti found G	Zr added G	Zr found G
1	0.0005	0.0009	0.0034	0.0009	None	None	None	None	None	None
2	.0005	.0009	. 0034	. 0009	None	None	None	None	None	None
3	. 0005	.0009	. 0034	.0009	0.0100	0.0101	0.0100	a 0. 0102	0.0101	b 0.0097
4	.0005	. 0009	. 0034	.0009	.0100	. 0094	.0100	a. 0102	.0101	b. 0097
5	. 0005	.0009	.0034	.0009	. 0500	.0502	. 0476	c. 0482	. 0500	b. 0493
6	.0005	. 0009	.0004	.0009	. 0500	. 0501	. 0476	c. 0482	. 0500	b. 0492
	b									

a Colorimetrically.

b The special treatment for vanadium (see note 2) was not carried out. This furnishes an interesting light on the slightly higher values for titanium obtained both colorimetrically and volumetrically and the correspondingly lower values for zirconium which resulted on account of the omission of this step.

c Volumetrically after reduction in a Jones reductor and collection in ferric-alum solution.

Loc. cit.

⁴ Technology Quarterly, 21, p. 35, 1908.

⁶ Am. J. Sci., 87, p. 173, 1914.

The following modifications of the above method were employed by Lieut. R. McLane in the analysis of zirconium steels at the Ithaca station of the Bureau of Mines:

1. Treat the evaporated solution containing silica with 25 cc of hydrochloric acid (sp. gr. 1.2), again evaporate to dryness, bake and take up in 30 cc of hydrochloric

acid (sp. gr. 1.2)+40 cc of water.

- 2. Ignite the insoluble residue and without weighing (silica being obtained on a separate sample by dehydration with sulphuric acid) add 2 cc of sulphuric acid (sp. gr. 1.84), an excess of hydrofluoric acid, and fume off the sulphuric acid. Dissolve the unignited residue in hydrochloric acid (1:1) and add to the acid extract from the ether separation.
- 3. Evaporate the filtrate from the silica determination to 25-40 cc volume, cool by placing in a larger beaker through which a stream of water is passed, and add 200 cc of ether. Stir, let settle, decant off ether, add 100 cc more ether, and repeat the operation. Perform a third extraction, if necessary, and pipette off the last of the ether, thus avoiding any transfer of the solution.
- 4. To separate aluminum, heat the oxidized solution to boiling and pour it with constant stirring into 135 cc of hot sodium hydroxide solution (20 per cent) contained in a 600 cc Pyrex beaker. After the precipitate has settled filter, allow the filtrate to stand overnight, and refilter if a precipitate appears. One extraction carried on as above is sufficient.
- 5. Place the filters and precipitates in the original beaker, add 25 cc of hydrochloric acid (sp. gr. 1.2), dilute to 125 cc, and heat. Filter off the insoluble, wash, ignite, fuse with sodium carbonate, and proceed as in the method.
- 6. Add methyl red and 8 cc of ammonia (sp. gr. o.9) to the sodium-hydroxide filtrate, make slightly acid with hydrochloric acid, dilute to 500 cc, heat to boiling, and make just alkaline with ammonia. Let stand warm for one hour, filter, dissolve the precipitate in hydrochloric acid, and dilute the cooled solution to 100 cc volume. Take out exactly 10 cc for a Fe₂O₃ determination by the colorimetric thiocyanate method. Precipitate the remainder of the solution as above and proceed with the molybdate separation as in the method. Finally deduct nine-tenths of the blank (blank has had SiO₂ and Fe₂O₃ deducted and is usually negligible) and divide the weight of Al₂O₃ by 0.9, giving Al₂O₃.

7. Treat the ignited cupferron precipitate with an excess of sulphuric and hydrofluoric acids, evaporate, ignite, and weigh in order to correct for any silica present.

8. Dissolve the weighed precipitate in sulphuric and hydrofluoric acids, evaporate to fumes of sulphuric acid, and make up to definite volume. Determine TiO₂ in one aliquot portion by the colorimetric peroxide method and Fe₂O₃ in another by the colorimetric thiocyanate method and deduct.

The authors desire to express their thanks to Dr. W. F. Hillebrand for valuable suggestions and advice.

